## ZYVOX®

(linezolid) injection (linezolid) tablets (linezolid) for oral suspension

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To reduce the development of drug-resistant bacteria and maintain the effectiveness of ZYVOX formulations and other antibacterial drugs, ZYVOX should be used only to treat or prevent infections that are proven or strongly suspected to be caused by bacteria.

## **DESCRIPTION**

ZYVOX I.V. Injection, ZYVOX Tablets, and ZYVOX for Oral Suspension contain linezolid, which is a synthetic antibacterial agent of the oxazolidinone class. The chemical name for linezolid is (S)-N-[[3-[3-Fluoro-4-(4-morpholinyl)phenyl]-2-oxo-5-oxazolidinyl] methyl]-acetamide.

The empirical formula is  $C_{16}H_{20}FN_3O_4$ . Its molecular weight is 337.35, and its chemical structure is represented below:

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O & O & O \\
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ZYVOX I.V. Injection is supplied as a ready-to-use sterile isotonic solution for intravenous infusion. Each mL contains 2 mg of linezolid. Inactive ingredients are sodium citrate, citric acid, and dextrose in an aqueous vehicle for intravenous administration. The sodium (Na<sup>+</sup>) content is 0.38 mg/mL (5 mEq per 300-mL bag; 3.3 mEq per 200-mL bag; and 1.7 mEq per 100-mL bag).

ZYVOX Tablets for oral administration contain 400 mg or 600 mg linezolid as film-coated compressed tablets. Inactive ingredients are corn starch, microcrystalline cellulose, hydroxypropylcellulose, sodium starch glycolate, magnesium stearate, hypromellose, polyethylene glycol, titanium dioxide, and carnauba wax. The sodium (Na<sup>+</sup>) content is 1.95 mg per 400-mg tablet and 2.92 mg per 600-mg tablet (0.1 mEq per tablet, regardless of strength).

ZYVOX for Oral Suspension is supplied as an orange-flavored granule/powder for constitution into a suspension for oral administration. Following constitution, each 5 mL contains 100 mg of linezolid. Inactive ingredients are sucrose, citric acid, sodium citrate, microcrystalline cellulose and carboxymethylcellulose sodium, aspartame, xanthan gum, mannitol, sodium benzoate, colloidal silicon dioxide, sodium chloride, and flavors (see **PRECAUTIONS, Information for Patients**). The sodium (Na<sup>+</sup>) content is 8.52 mg per 5 mL (0.4 mEq per 5 mL).

## **CLINICAL PHARMACOLOGY**

## **Pharmacodynamics**

In a randomized, positive- and placebo-controlled crossover thorough QT study, 40 healthy subjects were administered a single ZYVOX 600 mg dose via a 1 hour IV infusion, a single ZYVOX 1200 mg dose via a 1 hour IV infusion, placebo, and a single oral dose of positive control. At both the 600 mg and 1200 mg ZYVOX doses, no significant effect on QTc interval was detected at peak plasma concentration or at any other time.

#### **Pharmacokinetics**

The mean pharmacokinetic parameters of linezolid in adults after single and multiple oral and intravenous (IV) doses are summarized in Table 1. Plasma concentrations of linezolid at steady-state after oral doses of 600 mg given every 12 hours (q12h) are shown in Figure 1.

Table 1. Mean (Standard Deviation) Pharmacokinetic Parameters of Linezolid in Adults

	$C_{max}$	$C_{min}$	T <sub>max</sub>	AUC *	t <sub>1/2</sub>	CL
<b>Dose of Linezolid</b>	μg/mL	μg/mL	hrs	μg • h/mL	hrs	mL/min
400 mg tablet						
single dose †	8.10		1.52	55.10	5.20	146
	(1.83)		(1.01)	(25.00)	(1.50)	(67)
every 12 hours	11.00	3.08	1.12	73.40	4.69	110
,	(4.37)	(2.25)	(0.47)	(33.50)	(1.70)	(49)
600 mg tablet						
single dose	12.70		1.28	91.40	4.26	127
-	(3.96)		(0.66)	(39.30)	(1.65)	(48)
every 12 hours	21.20	6.15	1.03	138.00	5.40	80
,	(5.78)	(2.94)	(0.62)	(42.10)	(2.06)	(29)
600 mg IV injection ‡						
single dose	12.90		0.50	80.20	4.40	138
-	(1.60)		(0.10)	(33.30)	(2.40)	(39)
every 12 hours	15.10	3.68	0.51	89.70	4.80	123
<u> </u>	(2.52)	(2.36)	(0.03)	(31.00)	(1.70)	(40)
600 mg oral						
suspension	11.00		0.97	80.80	4.60	141
single dose	(2.76)		(0.88)	(35.10)	(1.71)	(45)

<sup>\*</sup> AUC for single dose =  $AUC_{0-\infty}$ ; for multiple-dose =  $AUC_{0-\tau}$ 

<sup>†</sup> Data dose-normalized from 375 mg

Data dose-normalized from 625 mg, IV dose was given as 0.5-hour infusion.

 $C_{max}$  = Maximum plasma concentration;  $C_{min}$  = Minimum plasma concentration;  $T_{max}$  = Time to  $C_{max}$ ; AUC =



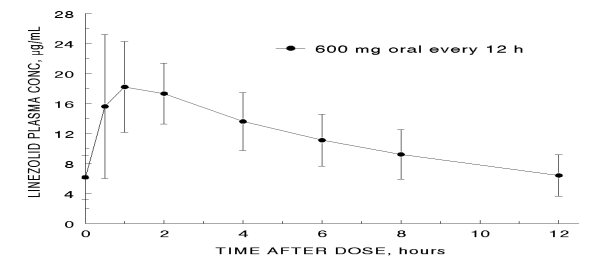


Figure 1. Plasma Concentrations of Linezolid in Adults at Steady-State Following Oral Dosing Every 12 Hours (Mean ± Standard Deviation, n=16)

**Absorption:** Linezolid is rapidly and extensively absorbed after oral dosing. Maximum plasma concentrations are reached approximately 1 to 2 hours after dosing, and the absolute bioavailability is approximately 100%. Therefore, linezolid may be given orally or intravenously without dose adjustment.

Linezolid may be administered without regard to the timing of meals. The time to reach the maximum concentration is delayed from 1.5 hours to 2.2 hours and  $C_{max}$  is decreased by about 17% when high fat food is given with linezolid. However, the total exposure measured as  $AUC_{0-\infty}$  values is similar under both conditions.

**Distribution:** Animal and human pharmacokinetic studies have demonstrated that linezolid readily distributes to well-perfused tissues. The plasma protein binding of linezolid is approximately 31% and is concentration-independent. The volume of distribution of linezolid at steady-state averaged 40 to 50 liters in healthy adult volunteers.

Linezolid concentrations have been determined in various fluids from a limited number of subjects in Phase 1 volunteer studies following multiple dosing of linezolid. The ratio of linezolid in saliva relative to plasma was 1.2 to 1 and for sweat relative to plasma was 0.55 to 1.

*Metabolism:* Linezolid is primarily metabolized by oxidation of the morpholine ring, which results in two inactive ring-opened carboxylic acid metabolites: the aminoethoxyacetic acid metabolite (A), and the hydroxyethyl glycine metabolite (B). Formation of metabolite A is presumed to be formed via an enzymatic pathway whereas metabolite B is mediated by a non-enzymatic chemical oxidation mechanism in vitro. In vitro studies have demonstrated

that linezolid is minimally metabolized and may be mediated by human cytochrome P450. However, the metabolic pathway of linezolid is not fully understood.

*Excretion:* Nonrenal clearance accounts for approximately 65% of the total clearance of linezolid. Under steady-state conditions, approximately 30% of the dose appears in the urine as linezolid, 40% as metabolite B, and 10% as metabolite A. The renal clearance of linezolid is low (average 40 mL/min) and suggests net tubular reabsorption. Virtually no linezolid appears in the feces, while approximately 6% of the dose appears in the feces as metabolite B, and 3% as metabolite A.

A small degree of nonlinearity in clearance was observed with increasing doses of linezolid, which appears to be due to lower renal and nonrenal clearance of linezolid at higher concentrations. However, the difference in clearance was small and was not reflected in the apparent elimination half-life.

## **Special Populations**

*Geriatric:* The pharmacokinetics of linezolid are not significantly altered in elderly patients (65 years or older). Therefore, dose adjustment for geriatric patients is not necessary.

**Pediatric:** The pharmacokinetics of linezolid following a single IV dose were investigated in pediatric patients ranging in age from birth through 17 years (including premature and full-term neonates), in healthy adolescent subjects ranging in age from 12 through 17 years, and in pediatric patients ranging in age from 1 week through 12 years. The pharmacokinetic parameters of linezolid are summarized in Table 2 for the pediatric populations studied and healthy adult subjects after administration of single IV doses.

The  $C_{max}$  and the volume of distribution ( $V_{ss}$ ) of linezolid are similar regardless of age in pediatric patients. However, clearance of linezolid varies as a function of age. With the exclusion of pre-term neonates less than one week of age, clearance is most rapid in the youngest age groups ranging from >1 week old to 11 years, resulting in lower single-dose systemic exposure (AUC) and shorter half-life as compared with adults. As age of pediatric patients increases, the clearance of linezolid gradually decreases, and by adolescence mean clearance values approach those observed for the adult population. There is wider intersubject variability in linezolid clearance and systemic drug exposure (AUC) across all pediatric age groups as compared with adults.

Similar mean daily AUC values were observed in pediatric patients from birth to 11 years of age dosed every 8 hours (q8h) relative to adolescents or adults dosed every 12 hours (q12h). Therefore, the dosage for pediatric patients up to 11 years of age should be 10 mg/kg q8h. Pediatric patients 12 years and older should receive 600 mg q12h (see **DOSAGE AND ADMINISTRATION**).

Table 2. Pharmacokinetic Parameters of Linezolid in Pediatrics and Adults Following a Single Intravenous Infusion of 10 mg/kg or 600 mg Linezolid (Mean: (%CV); [Min, Max Values])

Age Group	С <sub>тах</sub>	V <sub>ss</sub>	AUC <sup>*</sup>	t <sub>1/2</sub>	CL
	µg/mL	L/kg	μg•h/mL	hrs	mL/min/kg
Neonatal Patients Pre-term** < 1 week (N=9)†	12.7 (30%)	0.81 (24%)	108 (47%)	5.6 (46%)	2.0 (52%)
	[9.6, 22.2]	[0.43, 1.05]	[41, 191]	[2.4, 9.8]	[0.9, 4.0]
Full-term*** $< 1 \text{ week } (N=10)^{\dagger}$	11.5 (24%)	0.78 (20%)	55 (47%)	3.0 (55%)	3.8 (55%)
	[8.0, 18.3]	[0.45, 0.96]	[19, 103]	[1.3, 6.1]	[1.5, 8.8]
Full-term*** $\geq 1$ week to $\leq 28$ days $(N=10)^{\dagger}$	12.9 (28%)	0.66 (29%)	34 (21%)	1.5 (17%)	5.1 (22%)
	[7.7, 21.6]	[0.35, 1.06]	[23, 50]	[1.2, 1.9]	[3.3, 7.2]
Infant Patients > 28 days to < 3 Months (N=12) <sup>†</sup>	11.0 (27%)	0.79 (26%)	33 (26%)	1.8 (28%)	5.4 (32%)
	[7.2, 18.0]	[0.42, 1.08]	[17, 48]	[1.2, 2.8]	[3.5, 9.9]
Pediatric Patients 3 months through 11 years <sup>†</sup> (N=59)	15.1 (30%)	0.69 (28%)	58 (54%)	2.9 (53%)	3.8 (53%)
	[6.8, 36.7]	[0.31, 1.50]	[19, 153]	[0.9, 8.0]	[1.0, 8.5]
Adolescent Subjects and Patients 12 through 17 years <sup>‡</sup> (N=36)	16.7 (24%)	0.61 (15%)	95 (44%)	4.1 (46%)	2.1 (53%)
	[9.9, 28.9]	[0.44, 0.79]	[32, 178]	[1.3, 8.1]	[0.9, 5.2]
Adult Subjects <sup>§</sup> (N= 29)	12.5 (21%)	0.65 (16%)	91 (33%)	4.9 (35%)	1.7 (34%)
	[8.2, 19.3]	[0.45, 0.84]	[53, 155]	[1.8, 8.3]	[0.9, 3.3]

 $<sup>\</sup>overline{AUC}$  = Single dose  $AUC_{0-\infty}$ 

 $C_{max}$  = Maximum plasma concentration;  $V_{ss=}$  Volume of distribution; AUC = Area under concentration-time curve;

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*Gender:* Females have a slightly lower volume of distribution of linezolid than males. Plasma concentrations are higher in females than in males, which is partly due to body weight differences. After a 600-mg dose, mean oral clearance is approximately 38% lower in females than in males. However, there are no significant gender differences in mean apparent elimination-rate constant or half-life. Thus, drug exposure in females is not expected to substantially increase beyond levels known to be well tolerated. Therefore, dose adjustment by gender does not appear to be necessary.

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**Renal Insufficiency:** The pharmacokinetics of the parent drug, linezolid, are not altered in patients with any degree of renal insufficiency; however, the two primary metabolites of linezolid may accumulate in patients with renal insufficiency, with the amount of accumulation increasing with the severity of renal dysfunction (see Table 3). The clinical significance of accumulation of these two metabolites has not been determined in patients

In this data set, "pre-term" is defined as <34 weeks gestational age (Note: Only 1 patient enrolled was pre-term with a postnatal age between 1 week and 28 days)

<sup>\*\*\*</sup> In this data set, "full-term" is defined as ≥34 weeks gestational age

<sup>†</sup> Dose of 10 mg/kg

Dose of 600 mg or 10 mg/kg up to a maximum of 600 mg

<sup>§</sup> Dose normalized to 600 mg

 $t_{1/2}$  = Apparent elimination half-life; CL = Systemic clearance normalized for body weight

with severe renal insufficiency. Because similar plasma concentrations of linezolid are 146 147 achieved regardless of renal function, no dose adjustment is recommended for patients with renal insufficiency. However, given the absence of information on the clinical significance 148 149 of accumulation of the primary metabolites, use of linezolid in patients with renal 150 insufficiency should be weighed against the potential risks of accumulation of these metabolites. Both linezolid and the two metabolites are eliminated by dialysis. No 151 information is available on the effect of peritoneal dialysis on the pharmacokinetics of 152 linezolid. Approximately 30% of a dose was eliminated in a 3-hour dialysis session 153 154 beginning 3 hours after the dose of linezolid was administered; therefore, linezolid should 155 be given after hemodialysis. 156

Table 3. Mean (Standard Deviation) AUCs and Elimination Half-lives of Linezolid and Metabolites A and B in Patients with Varying Degrees of Renal Insufficiency After a Single 600-mg Oral Dose of Linezolid

Parameter	Healthy Subjects	Moderate Renal Impairment	Severe Renal Impairment	Hemodialysis-Dependent		
	$CL_{CR} > 80$ mL/min	$30 < CL_{CR} < 80 \text{ mL/min}$	$10 < CL_{CR} < 30 \text{ mL/min}$	Off Dialysis*	On Dialysis	
Linezolid						
AUC <sub>0-∞</sub> , μg h/mL	110 (22)	128 (53)	127 (66)	141 (45)	83 (23)	
t <sub>1/2</sub> , hours	6.4 (2.2)	6.1 (1.7)	7.1 (3.7)	8.4 (2.7)	7.0 (1.8)	
		Metab	olite A			
AUC <sub>0-48</sub> , μg h/mL	7.6 (1.9)	11.7 (4.3)	56.5 (30.6)	185 (124)	68.8 (23.9)	
t <sub>1/2</sub> , hours	6.3 (2.1)	6.6 (2.3)	9.0 (4.6)	NA	NA	
Metabolite B						
AUC <sub>0-48</sub> , μg h/mL	30.5 (6.2)	51.1 (38.5)	203 (92)	467 (102)	239 (44)	
t <sub>1/2</sub> , hours	6.6 (2.7)	9.9 (7.4)	11.0 (3.9)	NA	NA	

between hemodialysis sessions

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*Hepatic Insufficiency:* The pharmacokinetics of linezolid are not altered in patients (n=7) with mild-to-moderate hepatic insufficiency (Child-Pugh class A or B). On the basis of the available information, no dose adjustment is recommended for patients with mild-tomoderate hepatic insufficiency. The pharmacokinetics of linezolid in patients with severe hepatic insufficiency have not been evaluated.

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## **Drug-Drug Interactions**

Drugs Metabolized by Cytochrome P450: Linezolid is not an inducer of cytochrome P450 (CYP450) in rats. In addition, linezolid does not inhibit the activities of clinically significant human CYP isoforms (e.g., 1A2, 2C9, 2C19, 2D6, 2E1, 3A4). Therefore, linezolid is not expected to affect the pharmacokinetics of other drugs metabolized by these major enzymes. Concurrent administration of linezolid does not substantially alter the pharmacokinetic characteristics of (S)-warfarin, which is extensively metabolized by CYP2C9. Drugs such as warfarin and phenytoin, which are CYP2C9 substrates, may be given with linezolid without changes in dosage regimen.

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#### Antibiotics:

Aztreonam: The pharmacokinetics of linezolid or aztreonam are not altered when 175

administered together. 176

NA = Not applicable

*Gentamicin:* The pharmacokinetics of linezolid or gentamicin are not altered when administered together.

*Rifampin*: The effect of rifampin on the pharmacokinetics of linezolid was evaluated in a study of 16 healthy adult males. Volunteers were administered oral linezolid 600 mg twice daily for 5 doses with and without rifampin 600 mg once daily for 8 days. Coadministration of rifampin with linezolid resulted in a 21% decrease in linezolid  $C_{max}$  [90% CI, 15% - 27%] and a 32% decrease in linezolid  $AUC_{0-12}$  [90% CI, 27% - 37%]. The mechanism of this interaction is not fully understood and may be related to the induction of hepatic enzymes (see **PRECAUTIONS**, **Drug Interactions**).

*Monoamine Oxidase Inhibition:* Linezolid is a reversible, nonselective inhibitor of monoamine oxidase. Therefore, linezolid has the potential for interaction with adrenergic and serotonergic agents.

 Adrenergic Agents: A significant pressor response has been observed in normal adult subjects receiving linezolid and tyramine doses of more than 100 mg. Therefore, patients receiving linezolid need to avoid consuming large amounts of foods or beverages with high tyramine content (see **PRECAUTIONS**, **Information for Patients**).

A reversible enhancement of the pressor response of either pseudoephedrine HCl (PSE) or phenylpropanolamine HCl (PPA) is observed when linezolid is administered to healthy normotensive subjects (see PRECAUTIONS, Drug Interactions). A similar study has not been conducted in hypertensive patients. The interaction studies conducted in normotensive subjects evaluated the blood pressure and heart rate effects of placebo, PPA or PSE alone, linezolid alone, and the combination of steady-state linezolid (600 mg q12h for 3 days) with two doses of PPA (25 mg) or PSE (60 mg) given 4 hours apart. Heart rate was not affected by any of the treatments. Blood pressure was increased with both combination treatments. Maximum blood pressure levels were seen 2 to 3 hours after the second dose of PPA or PSE, and returned to baseline 2 to 3 hours after peak. The results of the PPA study follow, showing the mean (and range) maximum systolic blood pressure in mm Hg: placebo = 121 (103 to 158); linezolid alone = 120 (107 to 135); PPA alone = 125 (106 to 139); PPA with linezolid = 147 (129 to 176). The results from the PSE study were similar to those in the PPA study. The mean maximum increase in systolic blood pressure over baseline was 32 mm Hg (range: 20-52 mm Hg) and 38 mm Hg (range: 18-79 mm Hg) during co-administration of linezolid with pseudoephedrine or phenylpropanolamine, respectively.

Serotonergic Agents: The potential drug-drug interaction with dextromethorphan was
 studied in healthy volunteers. Subjects were administered dextromethorphan (two 20-mg
 doses given 4 hours apart) with or without linezolid. No serotonin syndrome effects
 (confusion, delirium, restlessness, tremors, blushing, diaphoresis, hyperpyrexia) have been
 observed in normal subjects receiving linezolid and dextromethorphan.

#### MICROBIOLOGY

Linezolid is a synthetic antibacterial agent of a new class of antibiotics, the oxazolidinones, 222 which has clinical utility in the treatment of infections caused by aerobic Gram-positive 223 224 bacteria. The in vitro spectrum of activity of linezolid also includes certain Gram-negative 225 bacteria and anaerobic bacteria. Linezolid inhibits bacterial protein synthesis through a 226 mechanism of action different from that of other antibacterial agents; therefore, cross-227 resistance between linezolid and other classes of antibiotics is unlikely. Linezolid binds to a site on the bacterial 23S ribosomal RNA of the 50S subunit and prevents the formation of 228 229 a functional 70S initiation complex, which is an essential component of the bacterial 230 translation process. The results of time-kill studies have shown linezolid to be bacteriostatic 231 against enterococci and staphylococci. For streptococci, linezolid was found to be 232 bactericidal for the majority of strains.

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In clinical trials, resistance to linezolid developed in 6 patients infected with *Enterococcus* faecium (4 patients received 200 mg q12h, lower than the recommended dose, and 2 patients received 600 mg q12h). In a compassionate use program, resistance to linezolid developed in 8 patients with E. faecium and in 1 patient with Enterococcus faecalis. All patients had either unremoved prosthetic devices or undrained abscesses. Resistance to linezolid occurs in vitro at a frequency of 1 x  $10^{-9}$  to 1 x  $10^{-11}$ . In vitro studies have shown that point mutations in the 23S rRNA are associated with linezolid resistance. Reports of vancomycin-resistant E. faecium becoming resistant to linezolid during its clinical use have been published. In one report nosocomial spread of vancomycin- and linezolid-resistant E. faecium occurred <sup>2</sup>. There has been a report of Staphylococcus aureus (methicillin-resistant) developing resistance to linezolid during its clinical use.<sup>3</sup> The linezolid resistance in these organisms was associated with a point mutation in the 23S rRNA (substitution of thymine for guanine at position 2576) of the organism. When antibiotic-resistant organisms are encountered in the hospital, it is important to emphasize infection control policies.<sup>4,5</sup> Resistance to linezolid has not been reported in *Streptococcus* spp., including Streptococcus pneumoniae.

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In vitro studies have demonstrated additivity or indifference between linezolid and vancomycin, gentamicin, rifampin, imipenem-cilastatin, aztreonam, ampicillin, or streptomycin.

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Linezolid has been shown to be active against most isolates of the following microorganisms, both in vitro and in clinical infections, as described in the **INDICATIONS AND USAGE** section.

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259	Aerobic and facultative Gram-positive microorganisms
260	Enterococcus faecium (vancomycin-resistant strains only)
261	Staphylococcus aureus (including methicillin-resistant strains)
262	Streptococcus agalactiae
263	Streptococcus pneumoniae (including multi-drug resistant isolates [MDRSP]*)
264	Streptococcus pyogenes
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266	The following in vitro data are available, but their clinical significance is unknown. At
267	least 90% of the following microorganisms exhibit an in vitro minimum inhibitory
268	concentration (MIC) less than or equal to the susceptible breakpoint for linezolid.
269	However, the safety and effectiveness of linezolid in treating clinical infections due to these
270	microorganisms have not been established in adequate and well-controlled clinical trials.
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272	Aerobic and facultative Gram-positive microorganisms
273	Enterococcus faecalis (including vancomycin-resistant strains)
274	Enterococcus faecium (vancomycin-susceptible strains)
275	Staphylococcus epidermidis (including methicillin-resistant strains)
276	Staphylococcus haemolyticus
277	Viridans group streptococci
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279	Aerobic and facultative Gram-negative microorganisms
280	Pasteurella multocida
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282	Susceptibility Testing Methods
283	<b>NOTE:</b> Susceptibility testing by dilution methods requires the use of linezolid
284	susceptibility powder.
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286	When available, the results of in vitro susceptibility tests should be provided to the
287	physician as periodic reports which describe the susceptibility profile of nosocomial and
288	community-acquired pathogens. These reports should aid the physician in selecting the
289	most effective antimicrobial.
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291	<b>Dilution Techniques:</b> Quantitative methods are used to determine antimicrobial minimum
292	inhibitory concentrations (MICs). These MICs provide estimates of the susceptibility of
293	bacteria to antimicrobial compounds. The MICs should be determined using a standardized
294	procedure. Standardized procedures are based on a dilution method <sup>6,7</sup> (broth or agar) or
295	equivalent with standardized inoculum concentrations and standardized concentrations of
296	linezolid powder. The MIC values should be interpreted according to criteria provided in

300 301 Table 4.

*Diffusion Techniques:* Quantitative methods that require measurement of zone diameters also provide reproducible estimates of the susceptibility of bacteria to antimicrobial compounds. One such standardized procedure <sup>7,8</sup> requires the use of standardized inoculum

<sup>\*</sup> MDRSP refers to isolates resistant to two or more of the following antibiotics: penicillin, second-generation cephalosporins, macrolides, tetracycline, and trimethoprim/sulfamethoxazole.

concentrations. This procedure uses paper disks impregnated with 30  $\mu$ g of linezolid to test the susceptibility of microorganisms to linezolid. The disk diffusion interpretive criteria are provided in Table 4.

Table 4. Susceptibility Interpretive Criteria for Linezolid

	Susceptibility Interpretive Criteria						
Pathogen	Minimal Inhibitory Concentrations (MIC in µg/mL)			Disk Diffusion (Zone Diameters in mm)			
	S	I	R	S	I	R	
Enterococcus spp	≤ 2	4	≥8	≥ 23	21-22	≤20	
Staphylococcus spp a	≤4			≥ 21			
Streptococcus pneumoniae	≤2 <sup>b</sup>			≥ 21°			
Streptococcus spp other than S pneumoniae a	≤2 <sup>b</sup>			≥ 21°			

<sup>&</sup>lt;sup>a</sup> The current absence of data on resistant strains precludes defining any categories other than "Susceptible." Strains yielding test results suggestive of a "nonsusceptible" category should be retested, and if the result is confirmed, the isolate should be submitted to a reference laboratory for further testing.

A report of "Susceptible" indicates that the pathogen is likely to be inhibited if the antimicrobial compound in the blood reaches the concentrations usually achievable. A report of "Intermediate" indicates that the result should be considered equivocal, and, if the microorganism is not fully susceptible to alternative, clinically feasible drugs, the test should be repeated. This category implies possible clinical applicability in body sites where the drug is physiologically concentrated or in situations where high dosage of drug can be used. This category also provides a buffer zone which prevents small uncontrolled technical factors from causing major discrepancies in interpretation. A report of "Resistant" indicates that the pathogen is not likely to be inhibited if the antimicrobial compound in the blood reaches the concentrations usually achievable; other therapy should be selected.

## **Quality Control**

Standardized susceptibility test procedures require the use of quality control microorganisms to control the technical aspects of the test procedures. Standard linezolid powder should provide the following range of values noted in Table 5. **NOTE:** Quality control microorganisms are specific strains of organisms with intrinsic biological properties relating to resistance mechanisms and their genetic expression within bacteria; the specific strains used for microbiological quality control are not clinically significant.

b These interpretive standards for *S. pneumoniae* and *Streptococcus* spp. other than *S. pneumoniae* are applicable only to tests performed by broth microdilution using cationadjusted Mueller-Hinton broth with 2 to 5% lysed horse blood inoculated with a direct colony suspension and incubated in ambient air at 35°C for 20 to 24 hours.

These zone diameter interpretive standards are applicable only to tests performed using Mueller-Hinton agar supplemented with 5% defibrinated sheep blood inoculated with a direct colony suspension and incubated in 5% CO<sub>2</sub> at 35°C for 20 to 24 hours.

Table 5. Acceptable Quality Control Ranges for Linezolid to be Used in Validation of Susceptibility Test Results

	Acceptable Quality Control Ranges				
QC Strain	Minimum Inhibitory Concentration (MIC in µg/mL)	Disk Diffusion (Zone Diameters in mm)			
Enterococcus faecalis ATCC 29212	1 - 4	Not applicable			
Staphylococcus aureus ATCC 29213	1 - 4	Not applicable			
Staphylococcus aureus ATCC 25923	Not applicable	25 - 32			
Streptococcus pneumoniae ATCC 49619 <sup>d</sup>	0.50 - 2 <sup>e</sup>	25 - 34 <sup>f</sup>			

<sup>&</sup>lt;sup>d</sup> This organism may be used for validation of susceptibility test results when testing *Streptococcus* spp. other than *S. pneumoniae*.

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#### INDICATIONS AND USAGE

ZYVOX formulations are indicated in the treatment of the following infections caused by susceptible strains of the designated microorganisms (see PRECAUTIONS, Pediatric Use and DOSAGE AND ADMINISTRATION and CLINICAL STUDIES). Linezolid is not indicated for the treatment of Gram-negative infections. It is critical that specific Gram-negative therapy be initiated immediately if a concomitant Gram-negative pathogen is documented or suspected (see WARNINGS).

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Vancomycin-Resistant *Enterococcus faecium* infections, including cases with concurrent bacteremia (see CLINICAL STUDIES).

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**Nosocomial pneumonia** caused by *Staphylococcus aureus* (methicillin-susceptible and -resistant strains), or *Streptococcus pneumoniae* (including multi-drug resistant strains [MDRSP]).

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Complicated skin and skin structure infections, including diabetic foot infections, without concomitant osteomyelitis, caused by *Staphylococcus aureus* (methicillinsusceptible and -resistant strains), *Streptococcus pyogenes*, or *Streptococcus agalactiae*. ZYVOX has not been studied in the treatment of decubitus ulcers.

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Uncomplicated skin and skin structure infections caused by *Staphylococcus aureus* (methicillin-susceptible only) or *Streptococcus pyogenes*.

<sup>&</sup>lt;sup>e</sup> This quality control range for *S. pneumoniae* is applicable only to tests performed by broth microdilution using cation-adjusted Mueller-Hinton broth with 2 to 5% lysed horse blood inoculated with a direct colony suspension and incubated in ambient air at 35°C for 20 to 24 hours.

This quality control zone diameter range is applicable only to tests performed using Mueller-Hinton agar supplemented with 5% defibrinated sheep blood inoculated with a direct colony suspension and incubated in 5% CO<sub>2</sub> at 35°C for 20 to 24 hours.

- 352 Community-acquired pneumonia caused by Streptococcus pneumoniae (including multi-
- drug resistant strains [MDRSP]\*), including cases with concurrent bacteremia, or 353
- Staphylococcus aureus (methicillin-susceptible strains only). 354

- 356 To reduce the development of drug-resistant bacteria and maintain the effectiveness of
- ZYVOX and other antibacterial drugs, ZYVOX should be used only to treat or prevent 357
- 358 infections that are proven or strongly suspected to be caused by susceptible bacteria. When
- culture and susceptibility information are available, they should be considered in selecting 359
- 360 or modifying antibacterial therapy. In the absence of such data, local epidemiology and
- susceptibility patterns may contribute to the empiric selection of therapy. 361

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#### CONTRAINDICATIONS

- ZYVOX formulations are contraindicated for use in patients who have known
- hypersensitivity to linezolid or any of the other product components. 365

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#### 367 Monoamine Oxidase Inhibitors

- Linezolid should not be used in patients taking any medicinal product which inhibits 368
- monoamine oxidases A or B (e.g., phenelzine, isocarboxazid) or within two weeks of taking 369
- 370 any such medicinal product.

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## Potential Interactions Producing Elevation of Blood Pressure

- 373 Unless patients are monitored for potential increases in blood pressure, linezolid should not
- 374 be administered to patients with uncontrolled hypertension, pheochromocytoma,
- thyrotoxicosis and/or patients taking any of the following types of medications: directly and 375
- 376 indirectly acting sympathomimetic agents (e.g., pseudoephedrine), vasopressive agents
- (e.g., epinephrine, norepinephrine), dopaminergic agents (e.g., dopamine, dobutamine) (see 377
- 378 PRECAUTIONS, Drug Interactions).

379 380

## **Potential Serotonergic Interactions**

- 381 Unless patients are carefully observed for signs and/or symptoms of serotonin syndrome,
- 382 linezolid should not be administered to patients with carcinoid syndrome and/or patients
- taking any of the following medications: serotonin re-uptake inhibitors, tricvelic 383
- 384 antidepressants, serotonin 5-HT1 receptor agonists (triptans), meperidine or buspirone (see
- PRECAUTIONS, General and Drug Interactions). 385

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#### WARNINGS

- 388 Myelosuppression (including anemia, leukopenia, pancytopenia, and
- thrombocytopenia) has been reported in patients receiving linezolid. In cases where 389
- 390 the outcome is known, when linezolid was discontinued, the affected hematologic
- 391 parameters have risen toward pretreatment levels. Complete blood counts should be
- monitored weekly in patients who receive linezolid, particularly in those who receive 392
- linezolid for longer than two weeks, those with pre-existing myelosuppression, those 393

<sup>\*</sup>MDRSP refers to isolates resistant to two or more of the following antibiotics: penicillin, second-generation cephalosporins, macrolides, tetracycline, and trimethoprim/sulfamethoxazole.

receiving concomitant drugs that produce bone marrow suppression, or those with a chronic infection who have received previous or concomitant antibiotic therapy.

Discontinuation of therapy with linezolid should be considered in patients who develop or have worsening myelosuppression.

In adult and juvenile dogs and rats, myelosuppression, reduced extramedullary hematopoiesis in spleen and liver, and lymphoid depletion of thymus, lymph nodes, and spleen were observed (see **ANIMAL PHARMACOLOGY**).

# Mortality Imbalance in an Investigational Study in Patients with Catheter-Related Bloodstream Infections, including those with catheter-site infections

 An imbalance in mortality was seen in patients treated with linezolid relative to vancomycin/dicloxacillin/oxacillin in an open-label study in seriously ill patients with intravascular catheter-related infections [78/363 (21.5%) vs. 58/363 (16.0%); odds ratio 1.426, 95% CI 0.970, 2.098]. While causality has not been established, this observed imbalance occurred primarily in linezolid-treated patients in whom either Gram-negative pathogens, mixed Gram-negative and Gram-positive pathogens, or no pathogen were identified at baseline, but was not seen in patients with Gram-positive infections only.

Linezolid is not approved and should not be used for the treatment of patients with catheter-related bloodstream infections or catheter-site infections.

Linezolid has no clinical activity against Gram-negative pathogens and is not indicated for the treatment of Gram-negative infections. It is critical that specific Gram-negative therapy be initiated immediately if a concomitant Gram-negative pathogen is documented or suspected (see **INDICATIONS AND USAGE**).

Clostridium difficile associated diarrhea (CDAD) has been reported with use of nearly all antibacterial agents, including ZYVOX, and may range in severity from mild diarrhea to fatal colitis. Treatment with antibacterial agents alters the normal flora of the colon leading to overgrowth of *C. difficile*.

*C. difficile* produces toxins A and B which contribute to the development of CDAD. Hypertoxin producing strains of *C. difficile* cause increased morbidity and mortality, as these infections can be refractory to antimicrobial therapy and may require colectomy. CDAD must be considered in all patients who present with diarrhea following antibiotic use.

Careful medical history is necessary since CDAD has been reported to occur over two months after the administration of antibacterial agents.

If CDAD is suspected or confirmed, ongoing antibiotic use not directed against *C. difficile* may need to be discontinued. Appropriate fluid and electrolyte management, protein supplementation, antibiotic treatment of *C. difficile*, and surgical evaluation should be instituted as clinically indicated.

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441	HV	nogiv	cemia

- 442 Postmarketing cases of symptomatic hypoglycemia have been reported in patients with diabetes
- 443 mellitus receiving insulin or oral hypoglycemic agents when treated with linezolid, a reversible,
- 444 <u>nonselective MAO inhibitor. Some MAO inhibitors have been associated with hypoglycemic</u>
- episodes in diabetic patients receiving insulin or hypoglycemic agents. While a causal relationship
- between linezolid and hypoglycemia has not been established, diabetic patients should be cautioned
- of potential hypoglycemic reactions when treated with linezolid.
- 448 If hypoglycemia occurs, a decrease in the dose of insulin or oral hypoglycemic agent, or
- discontinuation of oral hypoglycemic agent, insulin, or linezolid may be required.

#### 451 **PRECAUTIONS**

- 452 General
- 453 Lactic Acidosis
- Lactic acidosis has been reported with the use of ZYVOX. In reported cases, patients
- 455 experienced repeated episodes of nausea and vomiting. Patients who develop
- recurrent nausea or vomiting, unexplained acidosis, or a low bicarbonate level while
- receiving **ZYVOX** should receive immediate medical evaluation.

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- 459 **Serotonin Syndrome**
- 460 Spontaneous reports of serotonin syndrome associated with the co-administration of
- 2461 ZYVOX and serotonergic agents, including antidepressants such as selective serotonin
- reuptake inhibitors (SSRIs), have been reported (see PRECAUTIONS, Drug
- 463 Interactions).

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- 465 Where administration of ZYVOX and concomitant serotonergic agents is clinically
- appropriate, patients should be closely observed for signs and symptoms of serotonin
- 467 syndrome such as cognitive dysfunction, hyperpyrexia, hyperreflexia and
- 468 incoordination. If signs or symptoms occur physicians should consider discontinuation
- of either one or both agents. If the concomitant serotonergic agent is withdrawn,
- 470 discontinuation symptoms can be observed (see package insert of the specified
- agent(s) for a description of the associated discontinuation symptoms).

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## Peripheral and Optic Neuropathy

- 474 Peripheral and optic neuropathy have been reported in patients treated with ZYVOX,
- primarily those patients treated for longer than the maximum recommended duration of 28
- days. In cases of optic neuropathy that progressed to loss of vision, patients were treated for
- extended periods beyond the maximum recommended duration. Visual blurring has been
- 478 reported in some patients treated with ZYVOX for less than 28 days.

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- 480 If patients experience symptoms of visual impairment, such as changes in visual acuity,
- changes in color vision, blurred vision, or visual field defect, prompt ophthalmic evaluation
- 482 is recommended. Visual function should be monitored in all patients taking ZYVOX
- 483 for extended periods (≥ 3 months) and in all patients reporting new visual symptoms
- regardless of length of therapy with ZYVOX. If peripheral or optic neuropathy occurs,
- 485 the continued use of ZYVOX in these patients should be weighed against the potential
- 486 risks.

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#### Convulsions

Convulsions have been reported in patients when treated with linezolid. In some of these 490 cases, a history of seizures or risk factors for seizures was reported. 491

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The use of antibiotics may promote the overgrowth of nonsusceptible organisms. Should superinfection occur during therapy, appropriate measures should be taken.

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ZYVOX has not been studied in patients with uncontrolled hypertension. pheochromocytoma, carcinoid syndrome, or untreated hyperthyroidism.

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The safety and efficacy of ZYVOX formulations given for longer than 28 days have not been evaluated in controlled clinical trials.

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Prescribing ZYVOX in the absence of a proven or strongly suspected bacterial infection or a prophylactic indication is unlikely to provide benefit to the patient and increases the risk of the development of drug-resistant bacteria.

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## **Information for Patients**

Patients should be advised that:

- ZYVOX may be taken with or without food.
- They should inform their physician if they have a history of hypertension.
- Large quantities of foods or beverages with high tyramine content should be avoided 510 while taking ZYVOX. Quantities of tyramine consumed should be less than 100 mg 511 per meal. Foods high in tyramine content include those that may have undergone 512 protein changes by aging, fermentation, pickling, or smoking to improve flavor, such as 513 aged cheeses (0 to 15 mg tyramine per ounce); fermented or air-dried meats (0.1 to 514 515 8 mg tyramine per ounce); sauerkraut (8 mg tyramine per 8 ounces); soy sauce (5 mg tyramine per 1 teaspoon); tap beers (4 mg tyramine per 12 ounces); red wines 516 (0 to 6 mg tyramine per 8 ounces). The tyramine content of any protein-rich food may 517 be increased if stored for long periods or improperly refrigerated. <sup>9,10</sup> 518
- They should inform their physician if taking medications containing pseudoephedrine 519 HCl or phenylpropanolamine HCl, such as cold remedies and decongestants. 520
- They should inform their physician if taking serotonin re-uptake inhibitors or other 521 antidepressants. 522
  - Phenylketonurics: Each 5 mL of the 100 mg/5 mL ZYVOX for Oral Suspension contains 20 mg phenylalanine. The other ZYVOX formulations do not contain phenylalanine. Contact your physician or pharmacist.
  - They should inform their physician if they experience changes in vision.
  - They should inform their physician if they have a history of seizures.
- Diarrhea is a common problem caused by antibiotics, which usually ends when the antibiotic is discontinued. Sometimes after starting treatment with antibiotics, patients can develop watery and bloody stools (with or without stomach cramps and fever) even as late as two or more months after having taken the last dose of the antibiotic. If this 532 occurs, patients should contact their physician as soon as possible.

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Patients should be counseled that antibacterial drugs including ZYVOX should only be 534 535 used to treat bacterial infections. They do not treat viral infections (e.g., the common cold). When ZYVOX is prescribed to treat a bacterial infection, patients should be told that although it is common to feel better early in the course of therapy, the medication should be taken exactly as directed. Skipping doses or not completing the full course of therapy may (1) decrease the effectiveness of the immediate treatment and (2) increase the likelihood that bacteria will develop resistance and will not be treatable by ZYVOX or other antibacterial drugs in the future.

- 543 Drug Interactions (see also CLINICAL PHARMACOLOGY, Drug-Drug Interactions)
- 544 *Monoamine Oxidase Inhibition:* Linezolid is a reversible, nonselective inhibitor of
- monoamine oxidase. Therefore, linezolid has the potential for interaction with adrenergic

and serotonergic agents.

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Adrenergic Agents: Some individuals receiving ZYVOX may experience a reversible enhancement of the pressor response to indirect-acting sympathomimetic agents, vasopressor or dopaminergic agents. Commonly used drugs such as phenylpropanolamine and pseudoephedrine have been specifically studied. Initial doses of adrenergic agents, such as dopamine or epinephrine, should be reduced and titrated to achieve the desired response.

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Serotonergic Agents: Co-administration of linezolid and serotonergic agents was not associated with serotonin syndrome in Phase 1, 2 or 3 studies. Spontaneous reports of serotonin syndrome associated with co-administration of ZYVOX and serotonergic agents, including antidepressants such as selective serotonin reuptake inhibitors (SSRIs), have been reported. Patients who are treated with ZYVOX and concomitant serotonergic agents should be closely observed as described in the PRECAUTIONS, General Section.

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Strong CYP450 Inducers: In a study in healthy volunteers, co-administration of rifampin with oral linezolid resulted in a 21% decrease in linezolid C<sub>max</sub> and a 32% decrease in linezolid AUC<sub>0-12</sub>. The clinical significance of this interaction is unknown. Other strong inducers of hepatic enzymes (e.g. carbamazepine, phenytoin, phenobarbital) could cause a similar or smaller decrease in linezolid exposure (see CLINICAL PHARMACOLOGY,

567 **Drug-Drug Interactions**)

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## **Drug-Laboratory Test Interactions**

There are no reported drug-laboratory test interactions.

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## Carcinogenesis, Mutagenesis, Impairment of Fertility

Lifetime studies in animals have not been conducted to evaluate the carcinogenic potential of linezolid. Neither mutagenic nor clastogenic potential was found in a battery of tests including: assays for mutagenicity (Ames bacterial reversion and CHO cell mutation), an in vitro unscheduled DNA synthesis (UDS) assay, an in vitro chromosome aberration assay in human lymphocytes, and an in vivo mouse micronucleus assay.

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579 Linezolid did not affect the fertility or reproductive performance of adult female rats. It reversibly decreased fertility and reproductive performance in adult male rats when given at 580 581 doses  $\geq 50$  mg/kg/day, with exposures approximately equal to or greater than the expected 582 human exposure level (exposure comparisons are based on AUCs). The reversible fertility effects were mediated through altered spermatogenesis. Affected spermatids contained 583 584 abnormally formed and oriented mitochondria and were non-viable. Epithelial cell 585 hypertrophy and hyperplasia in the epididymis was observed in conjunction with decreased 586 fertility. Similar epididymal changes were not seen in dogs.

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In sexually mature male rats exposed to drug as juveniles, mildly decreased fertility was observed following treatment with linezolid through most of their period of sexual

development (50 mg/kg/day from days 7 to 36 of age, and 100 mg/kg/day from days 37 to 55 of age), with exposures up to 1.7-fold greater than mean AUCs observed in pediatric patients aged 3 months to 11 years. Decreased fertility was not observed with shorter treatment periods, corresponding to exposure in utero through the early neonatal period (gestation day 6 through postnatal day 5), neonatal exposure (postnatal days 5 to 21), or to juvenile exposure (postnatal days 22 to 35). Reversible reductions in sperm motility and altered sperm morphology were observed in rats treated from postnatal day 22 to 35.

## **Pregnancy**

**Teratogenic Effects. Pregnancy Category C:** Linezolid was not teratogenic in mice, rats, or rabbits at exposure levels 6.5-fold (in mice), equivalent to (in rats), or 0.06-fold (in rabbits) the expected human exposure level, based on AUCs. However, embryo and fetal toxicities were seen (see **Non-teratogenic Effects**). There are no adequate and well-controlled studies in pregnant women. ZYVOX should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.

## **Non-teratogenic Effects**

In mice, embryo and fetal toxicities were seen only at doses that caused maternal toxicity (clinical signs and reduced body weight gain). A dose of 450 mg/kg/day (6.5-fold the estimated human exposure level based on AUCs) correlated with increased postimplantational embryo death, including total litter loss, decreased fetal body weights, and an increased incidence of costal cartilage fusion.

In rats, mild fetal toxicity was observed at 15 and 50 mg/kg/day (exposure levels 0.22-fold to approximately equivalent to the estimated human exposure, respectively based on AUCs). The effects consisted of decreased fetal body weights and reduced ossification of sternebrae, a finding often seen in association with decreased fetal body weights. Slight maternal toxicity, in the form of reduced body weight gain, was seen at 50 mg/kg/day.

 In rabbits, reduced fetal body weight occurred only in the presence of maternal toxicity (clinical signs, reduced body weight gain and food consumption) when administered at a dose of 15 mg/kg/day (0.06-fold the estimated human exposure based on AUCs).

 When female rats were treated with 50 mg/kg/day (approximately equivalent to the estimated human exposure based on AUCs) of linezolid during pregnancy and lactation, survival of pups was decreased on postnatal days 1 to 4. Male and female pups permitted to mature to reproductive age, when mated, showed an increase in preimplantation loss.

#### **Nursing Mothers**

Linezolid and its metabolites are excreted in the milk of lactating rats. Concentrations in milk were similar to those in maternal plasma. It is not known whether linezolid is excreted in human milk. Because many drugs are excreted in human milk, caution should be exercised when ZYVOX is administered to a nursing woman.

#### Pediatric Use

The safety and effectiveness of ZYVOX for the treatment of pediatric patients with the following infections are supported by evidence from adequate and well-controlled studies

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adults, pharmacokinetic data in pediatric patients, and additional data from a comparator-controlled study of Gram-positive infections in pediatric patients ranging in age from birth through 11 years (see INDICATIONS AND USAGE and CLINICAL STUDIES):

- nosocomial pneumonia
- complicated skin and skin structure infections
- community-acquired pneumonia (also supported by evidence from an uncontrolled study in patients ranging in age from 8 months through 12 years)
- vancomycin-resistant *Enterococcus faecium* infections

The safety and effectiveness of ZYVOX for the treatment of pediatric patients with the following infection have been established in a comparator-controlled study in pediatric patients ranging in age from 5 through 17 years (see CLINICAL STUDIES):

• uncomplicated skin and skin structure infections caused by *Staphylococcus aureus* (methicillin-susceptible strains only) or *Streptococcus pyogenes* 

Pharmacokinetic information generated in pediatric patients with ventriculoperitoneal shunts showed variable cerebrospinal fluid (CSF) linezolid concentrations following single and multiple dosing of linezolid; therapeutic concentrations were not consistently achieved or maintained in the CSF. Therefore, the use of linezolid for the empiric treatment of pediatric patients with central nervous system infections is not recommended.

The  $C_{max}$  and the volume of distribution ( $V_{ss}$ ) of linezolid are similar regardless of age in pediatric patients. However, linezolid clearance is a function of age. Excluding neonates less than a week of age, clearance is most rapid in the youngest age groups ranging from >1 week old to 11 years, resulting in lower single-dose systemic exposure (AUC) and shorter half-life as compared with adults. As age of pediatric patients increases, the clearance of linezolid gradually decreases, and by adolescence, mean clearance values approach those observed for the adult population. There is wider inter-subject variability in linezolid clearance and in systemic drug exposure (AUC) across all pediatric age groups as compared with adults.

667 with adults

Similar mean daily AUC values were observed in pediatric patients from birth to 11 years of age dosed q8h relative to adolescents or adults dosed q12h. Therefore, the dosage for pediatric patients up to 11 years of age should be 10 mg/kg q8h. Pediatric patients 12 years and older should receive 600 mg q12h.

Recommendations for the dosage regimen for pre-term neonates less than 7 days of age (gestational age less than 34 weeks) are based on pharmacokinetic data from 9 pre-term neonates. Most of these pre-term neonates have lower systemic linezolid clearance values and larger AUC values than many full-term neonates and older infants. Therefore, these pre-term neonates should be initiated with a dosing regimen of 10 mg/kg q12h. Consideration may be given to the use of a 10 mg/kg q8h regimen in neonates with a suboptimal clinical response. All neonatal patients should receive 10 mg/kg q8h by 7 days of

680 life (see CLINICAL PHARMACOLOGY, Special Populations, Pediatric and

**DOSAGE AND ADMINISTRATION**).

- In limited clinical experience, 5 out of 6 (83%) pediatric patients with infections due to
- 684 Gram-positive pathogens with MICs of 4 µg/mL treated with ZYVOX had clinical cures.
- However, pediatric patients exhibit wider variability in linezolid clearance and systemic
- exposure (AUC) compared with adults. In pediatric patients with a sub-optimal clinical
- response, particularly those with pathogens with MIC of 4  $\mu$ g/mL, lower systemic
- exposure, site and severity of infection, and the underlying medical condition should be
- considered when assessing clinical response (see CLINICAL PHARMACOLOGY,
- 690 Special Populations, Pediatric and DOSAGE AND ADMINISTRATION).

## Geriatric Use

- Of the 2046 patients treated with ZYVOX in Phase 3 comparator-controlled clinical trials,
- 589 (29%) were 65 years or older and 253 (12%) were 75 years or older. No overall
- differences in safety or effectiveness were observed between these patients and younger
- 696 patients.

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## ANIMAL PHARMACOLOGY

- Target organs of linezolid toxicity were similar in juvenile and adult rats and dogs. Dose-
- and time-dependent myelosuppression, as evidenced by bone marrow
- hypocellularity/decreased hematopoiesis, decreased extramedullary hematopoiesis in spleen
- and liver, and decreased levels of circulating erythrocytes, leukocytes, and platelets have
- been seen in animal studies. Lymphoid depletion occurred in thymus, lymph nodes, and
- spleen. Generally, the lymphoid findings were associated with anorexia, weight loss, and
- suppression of body weight gain, which may have contributed to the observed effects.

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- In rats administered linezolid orally for 6 months, non-reversible, minimal to mild axonal
- degeneration of sciatic nerves was observed at 80 mg/kg/day; minimal degeneration of the sciatic nerve was also observed in 1 male at this dose level at a 3-month interim necropsy.
- Sensitive morphologic evaluation of perfusion-fixed tissues was conducted to investigate
- evidence of optic nerve degeneration. Minimal to moderate optic nerve degeneration was
- evident in 2 male rats after 6 months of dosing, but the direct relationship to drug was
- equivocal because of the acute nature of the finding and its asymmetrical distribution. The
- nerve degeneration observed was microscopically comparable to spontaneous unilateral
- optic nerve degeneration reported in aging rats and may be an exacerbation of common
- 516 background change.

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- These effects were observed at exposure levels that are comparable to those observed in
- some human subjects. The hematopoietic and lymphoid effects were reversible, although in
- some studies, reversal was incomplete within the duration of the recovery period.

#### ADVERSE REACTIONS

#### **Adult Patients**

The safety of ZYVOX formulations was evaluated in 2046 adult patients enrolled in seven Phase 3 comparator-controlled clinical trials, who were treated for up to 28 days. In these studies, 85% of the adverse events reported with ZYVOX were described as mild to moderate in intensity. Table 6 shows the incidence of adverse events reported in at least 2% of patients in these trials. The most common adverse events in patients treated with ZYVOX were diarrhea (incidence across studies: 2.8% to 11.0%), headache (incidence across studies: 0.5% to 11.3%), and nausea (incidence across studies: 3.4% to 9.6%).

Table 6. Incidence (%) of Adverse Events Reported in ≥2% of Adult Patients in Comparator-Controlled Clinical Trials with ZYVOX

Event	ZYVOX (n=2046)	All Comparators * (n=2001)
Diarrhea	8.3	6.3
Headache	6.5	5.5
Nausea	6.2	4.6
Vomiting	3.7	2.0
Insomnia	2.5	1.7
Constipation	2.2	2.1
Rash	2.0	2.2
Dizziness	2.0	1.9
Fever	1.6	2.1

<sup>\*</sup> Comparators included cefpodoxime proxetil 200 mg PO q12h; ceftriaxone 1 g IV q12h; clarithromycin 250 mg PO q12h; dicloxacillin 500 mg PO q6h; oxacillin 2 g IV q6h; vancomycin 1 g IV q12h.

Other adverse events reported in Phase 2 and Phase 3 studies included oral moniliasis, vaginal moniliasis, hypertension, dyspepsia, localized abdominal pain, pruritus, and tongue discoloration.

Table 7 shows the incidence of drug-related adverse events reported in at least 1% of adult patients in these trials by dose of ZYVOX.

Table 7. Incidence (%) of Drug-Related Adverse Events Occurring in >1% of Adult Patients Treated with ZYVOX in Comparator-Controlled Clinical Trials

		d Skin and Skin Infections	All Other	Indications
Adverse Event	ZYVOX 400 mg PO q12h (n=548)	Clarithromycin 250 mg PO q12h (n=537)	ZYVOX 600 mg q12h (n=1498)	All Other Comparators* (n=1464)
% of patients with 1 drug- related adverse event	25.4	19.6	20.4	14.3
% of patients discontinuing due to drug-related adverse events <sup>†</sup>	3.5	2.4	2.1	1.7
Diarrhea	5.3	4.8	4.0	2.7
Nausea	3.5	3.5	3.3	1.8
Headache	2.7	2.2	1.9	1.0
Taste alteration	1.8	2.0	0.9	0.2
Vaginal moniliasis	1.6	1.3	1.0	0.4
Fungal infection	1.5	0.2	0.1	< 0.1
Abnormal liver function tests	0.4	0	1.3	0.5
Vomiting	0.9	0.4	1.2	0.4
Tongue discoloration	1.1	0	0.2	0
Dizziness	1.1	1.5	0.4	0.3
Oral moniliasis	0.4	0	1.1	0.4

<sup>\*</sup> Comparators included cefpodoxime proxetil 200 mg PO q12h; ceftriaxone 1 g IV q12h; dicloxacillin 500 mg PO q6h; oxacillin 2 g IV q6h; vancomycin 1 g IV q12h.

## **Pediatric Patients**

The safety of ZYVOX formulations was evaluated in 215 pediatric patients ranging in age from birth through 11 years, and in 248 pediatric patients aged 5 through 17 years (146 of these 248 were age 5 through 11 and 102 were age 12 to 17). These patients were enrolled in two Phase 3 comparator-controlled clinical trials and were treated for up to 28 days. In these studies, 83% and 99%, respectively, of the adverse events reported with ZYVOX were described as mild to moderate in intensity. In the study of hospitalized pediatric patients (birth through 11 years) with Gram-positive infections, who were randomized 2 to 1 (linezolid:vancomycin), mortality was 6.0% (13/215) in the linezolid arm and 3.0% (3/101) in the vancomycin arm. However, given the severe underlying illness in the patient population, no causality could be established. Table 8 shows the incidence of adverse events reported in at least 2% of pediatric patients treated with ZYVOX in these trials.

The most commonly reported drug-related adverse events leading to discontinuation in patients treated with ZYVOX were nausea, headache, diarrhea, and vomiting.

Table 8. Incidence (%) of Adverse Events Reported in ≥2% of Pediatric Patients Treated with ZYVOX in Comparator-Controlled Clinical Trials

Event         ZYVOX (n=248)         Cefadroxil (n=251)         ZYVOX (n=215)         Vancomycin (n=101)           Fever         2.9         3.6         14.1         14.1           Diarrhea         7.8         8.0         10.8         12.1           Vomiting         2.9         6.4         9.4         9.1           Sepsis         0         0         8.0         7.1           Rash         1.6         1.2         7.0         15.2           Headache         6.5         4.0         0.9         0           Anemia         0         0         5.6         7.1           Thrombocytopenia         0         0         4.7         2.0           Upper respiratory infection         3.7         5.2         4.2         1.0           Nausea         3.7         3.2         1.9         0           Dyspnea         0         0         3.3         5.1           Reaction at site of injection or of vascular catheter         0         0         3.3         5.1           Trauma         3.3         4.8         2.8         2.0           Pharyngitis         2.9         1.6         0.5         1.0           C			l Skin and Skin Infections*	All Other l	ndications <sup>†</sup>
Diarrhea         7.8         8.0         10.8         12.1           Vomiting         2.9         6.4         9.4         9.1           Sepsis         0         0         8.0         7.1           Rash         1.6         1.2         7.0         15.2           Headache         6.5         4.0         0.9         0           Anemia         0         0         5.6         7.1           Thrombocytopenia         0         0         4.7         2.0           Upper respiratory infection         3.7         5.2         4.2         1.0           Nausea         3.7         3.2         1.9         0           Dyspnea         0         0         3.3         1.0           Reaction at site of injection or of vascular catheter         0         0         3.3         5.1           Trauma         3.3         4.8         2.8         2.0           Pharyngitis         2.9         1.6         0.5         1.0           Convulsion         0         0         2.8         2.0           Hypokalemia         0         0         2.8         2.0           Thrombocythemia         0 <t< th=""><th>Event</th><th></th><th></th><th></th><th></th></t<>	Event				
Vomiting         2.9         6.4         9.4         9.1           Sepsis         0         0         8.0         7.1           Rash         1.6         1.2         7.0         15.2           Headache         6.5         4.0         0.9         0           Anemia         0         0         5.6         7.1           Thrombocytopenia         0         0         4.7         2.0           Upper respiratory infection         3.7         5.2         4.2         1.0           Nausea         3.7         3.2         1.9         0           Dyspnea         0         0         3.3         1.0           Reaction at site of injection or of vascular catheter         0         0         3.3         5.1           Trauma         3.3         4.8         2.8         2.0           Pharyngitis         2.9         1.6         0.5         1.0           Convulsion         0         0         2.8         2.0           Hypokalemia         0         0         2.8         2.0           Thrombocythemia         0         0         2.8         2.0           Cough         2.4         4.0 </td <td>Fever</td> <td>2.9</td> <td>3.6</td> <td>14.1</td> <td>14.1</td>	Fever	2.9	3.6	14.1	14.1
Sepsis         0         0         8.0         7.1           Rash         1.6         1.2         7.0         15.2           Headache         6.5         4.0         0.9         0           Anemia         0         0         5.6         7.1           Thrombocytopenia         0         0         4.7         2.0           Upper respiratory infection         3.7         5.2         4.2         1.0           Nausea         3.7         3.2         1.9         0           Dyspnea         0         0         3.3         1.0           Reaction at site of injection or of vascular catheter         0         0         3.3         5.1           Trauma         3.3         4.8         2.8         2.0           Pharyngitis         2.9         1.6         0.5         1.0           Convulsion         0         0         2.8         2.0           Hypokalemia         0         0         2.8         2.0           Thrombocythemia         0         0         2.8         2.0           Thrombocythemia         0         0         2.8         2.0           Cough         2.4         4.	Diarrhea	7.8	8.0	10.8	12.1
Rash         1.6         1.2         7.0         15.2           Headache         6.5         4.0         0.9         0           Anemia         0         0         5.6         7.1           Thrombocytopenia         0         0         4.7         2.0           Upper respiratory infection         3.7         5.2         4.2         1.0           Nausea         3.7         3.2         1.9         0           Dyspnea         0         0         3.3         1.0           Reaction at site of injection or of vascular catheter         0         0         3.3         5.1           Trauma         3.3         4.8         2.8         2.0           Pharyngitis         2.9         1.6         0.5         1.0           Convulsion         0         0         2.8         2.0           Hypokalemia         0         0         2.8         2.0           Hypokalemia         0         0         2.8         2.0           Thrombocythemia         0         0         2.8         2.0           Cough         2.4         4.0         0.9         0           Generalized abdominal pain         2.4 </td <td>Vomiting</td> <td>2.9</td> <td>6.4</td> <td>9.4</td> <td>9.1</td>	Vomiting	2.9	6.4	9.4	9.1
Headache         6.5         4.0         0.9         0           Anemia         0         0         5.6         7.1           Thrombocytopenia         0         0         4.7         2.0           Upper respiratory infection         3.7         5.2         4.2         1.0           Nausea         3.7         3.2         1.9         0           Dyspnea         0         0         3.3         1.0           Reaction at site of injection or of vascular catheter         0         0         3.3         5.1           Trauma         3.3         4.8         2.8         2.0           Pharyngitis         2.9         1.6         0.5         1.0           Convulsion         0         0         2.8         2.0           Hypokalemia         0         0         2.8         2.0           Thrombocythemia         0         0         2.8         2.0           Thrombocythemia         0         0         2.8         2.0           Cough         2.4         4.0         0.9         0           Generalized abdominal pain         2.4         2.8         0.9         2.0           Localized abdominal pain	Sepsis	0	0	8.0	7.1
Anemia         0         0         5.6         7.1           Thrombocytopenia         0         0         4.7         2.0           Upper respiratory infection         3.7         5.2         4.2         1.0           Nausea         3.7         3.2         1.9         0           Dyspnea         0         0         3.3         1.0           Reaction at site of injection or of vascular catheter         0         0         3.3         5.1           Trauma         3.3         4.8         2.8         2.0           Pharyngitis         2.9         1.6         0.5         1.0           Convulsion         0         0         2.8         2.0           Hypokalemia         0         0         2.8         2.0           Thrombocythemia         0         0         2.8         2.0           Thrombocythemia         0         0         2.8         2.0           Cough         2.4         4.0         0.9         0           Generalized abdominal pain         2.4         2.8         0.5         1.0           Localized abdominal pain         2.4         2.8         0.5         1.0           Gastro	Rash	1.6	1.2	7.0	15.2
Thrombocytopenia         0         4.7         2.0           Upper respiratory infection         3.7         5.2         4.2         1.0           Nausea         3.7         3.2         1.9         0           Dyspnea         0         0         3.3         1.0           Reaction at site of injection or of vascular catheter         0         0         3.3         5.1           Trauma         3.3         4.8         2.8         2.0           Pharyngitis         2.9         1.6         0.5         1.0           Convulsion         0         0         2.8         2.0           Hypokalemia         0         0         2.8         2.0           Thrombocythemia         0         0         2.8         2.0           Thrombocythemia         0         0         2.8         2.0           Cough         2.4         4.0         0.9         0           Generalized abdominal pain         2.4         2.8         0.9         2.0           Localized abdominal pain         2.4         2.8         0.5         1.0           Apnea         0         0         2.3         1.0           Generalized edema	Headache	6.5	4.0	0.9	0
Upper respiratory infection         3.7         5.2         4.2         1.0           Nausea         3.7         3.2         1.9         0           Dyspnea         0         0         3.3         1.0           Reaction at site of injection or of vascular catheter         0         0         3.3         5.1           Trauma         3.3         4.8         2.8         2.0           Pharyngitis         2.9         1.6         0.5         1.0           Convulsion         0         0         2.8         2.0           Hypokalemia         0         0         2.8         2.0           Pneumonia         0         0         2.8         2.0           Thrombocythemia         0         0         2.8         2.0           Cough         2.4         4.0         0.9         0           Generalized abdominal pain         2.4         2.8         0.9         2.0           Localized abdominal pain         2.4         2.8         0.5         1.0           Apnea         0         0         2.3         2.0           Gastrointestinal bleeding         0         0         2.3         1.0           Loos	Anemia	0	0	5.6	7.1
Nausea         3.7         3.2         1.9         0           Dyspnea         0         0         3.3         1.0           Reaction at site of injection or of vascular catheter         0         0         3.3         5.1           Trauma         3.3         4.8         2.8         2.0           Pharyngitis         2.9         1.6         0.5         1.0           Convulsion         0         0         2.8         2.0           Hypokalemia         0         0         2.8         2.0           Pneumonia         0         0         2.8         2.0           Thrombocythemia         0         0         2.8         2.0           Cough         2.4         4.0         0.9         0           Generalized abdominal pain         2.4         2.8         0.9         2.0           Localized abdominal pain         2.4         2.8         0.5         1.0           Apnea         0         0         2.3         2.0           Gastrointestinal bleeding         0         0         2.3         1.0           Generalized edema         0         0         2.3         1.0           Loose stools	Thrombocytopenia	0	0	4.7	2.0
Dyspnea         0         0         3.3         1.0           Reaction at site of injection or of vascular catheter         0         0         3.3         5.1           Trauma         3.3         4.8         2.8         2.0           Pharyngitis         2.9         1.6         0.5         1.0           Convulsion         0         0         2.8         2.0           Hypokalemia         0         0         2.8         2.0           Pneumonia         0         0         2.8         2.0           Thrombocythemia         0         0         2.8         2.0           Cough         2.4         4.0         0.9         0           Generalized abdominal pain         2.4         2.8         0.9         2.0           Localized abdominal pain         2.4         2.8         0.5         1.0           Apnea         0         0         2.3         2.0           Gastrointestinal bleeding         0         0         2.3         1.0           Loose stools         1.6         0.8         2.3         3.0           Localized pain         2.0         1.6         0.9         0	Upper respiratory infection	3.7	5.2	4.2	1.0
Reaction at site of injection or of vascular catheter         0         0         3.3         5.1           Trauma         3.3         4.8         2.8         2.0           Pharyngitis         2.9         1.6         0.5         1.0           Convulsion         0         0         2.8         2.0           Hypokalemia         0         0         2.8         2.0           Pneumonia         0         0         2.8         2.0           Thrombocythemia         0         0         2.8         2.0           Cough         2.4         4.0         0.9         0           Generalized abdominal pain         2.4         2.8         0.9         2.0           Localized abdominal pain         2.4         2.8         0.5         1.0           Apnea         0         0         2.3         2.0           Gastrointestinal bleeding         0         0         2.3         1.0           Generalized edema         0         0         2.3         1.0           Loose stools         1.6         0.8         2.3         3.0           Localized pain         2.0         1.6         0.9         0	Nausea	3.7	3.2	1.9	0
of vascular catheter         3.3         4.8         2.8         2.0           Pharyngitis         2.9         1.6         0.5         1.0           Convulsion         0         0         2.8         2.0           Hypokalemia         0         0         2.8         3.0           Pneumonia         0         0         2.8         2.0           Thrombocythemia         0         0         2.8         2.0           Cough         2.4         4.0         0.9         0           Generalized abdominal pain         2.4         2.8         0.9         2.0           Localized abdominal pain         2.4         2.8         0.5         1.0           Apnea         0         0         2.3         2.0           Gastrointestinal bleeding         0         0         2.3         1.0           Generalized edema         0         0         2.3         1.0           Localized pain         2.0         1.6         0.9         0	Dyspnea	0	0	3.3	1.0
Pharyngitis         2.9         1.6         0.5         1.0           Convulsion         0         0         2.8         2.0           Hypokalemia         0         0         2.8         3.0           Pneumonia         0         0         2.8         2.0           Thrombocythemia         0         0         2.8         2.0           Cough         2.4         4.0         0.9         0           Generalized abdominal pain         2.4         2.8         0.9         2.0           Localized abdominal pain         2.4         2.8         0.5         1.0           Apnea         0         0         2.3         2.0           Gastrointestinal bleeding         0         0         2.3         1.0           Generalized edema         0         0         2.3         1.0           Loose stools         1.6         0.8         2.3         3.0           Localized pain         2.0         1.6         0.9         0		0	0	3.3	5.1
Convulsion         0         0         2.8         2.0           Hypokalemia         0         0         2.8         3.0           Pneumonia         0         0         2.8         2.0           Thrombocythemia         0         0         2.8         2.0           Cough         2.4         4.0         0.9         0           Generalized abdominal pain         2.4         2.8         0.9         2.0           Localized abdominal pain         2.4         2.8         0.5         1.0           Apnea         0         0         2.3         2.0           Gastrointestinal bleeding         0         0         2.3         1.0           Generalized edema         0         0         2.3         1.0           Localized pain         2.0         1.6         0.8         2.3         3.0	Trauma	3.3	4.8	2.8	2.0
Hypokalemia         0         0         2.8         3.0           Pneumonia         0         0         2.8         2.0           Thrombocythemia         0         0         2.8         2.0           Cough         2.4         4.0         0.9         0           Generalized abdominal pain         2.4         2.8         0.9         2.0           Localized abdominal pain         2.4         2.8         0.5         1.0           Apnea         0         0         2.3         2.0           Gastrointestinal bleeding         0         0         2.3         1.0           Generalized edema         0         0         2.3         1.0           Loose stools         1.6         0.8         2.3         3.0           Localized pain         2.0         1.6         0.9         0	Pharyngitis	2.9	1.6	0.5	1.0
Pneumonia         0         0         2.8         2.0           Thrombocythemia         0         0         2.8         2.0           Cough         2.4         4.0         0.9         0           Generalized abdominal pain         2.4         2.8         0.9         2.0           Localized abdominal pain         2.4         2.8         0.5         1.0           Apnea         0         0         2.3         2.0           Gastrointestinal bleeding         0         0         2.3         1.0           Generalized edema         0         0         2.3         1.0           Loose stools         1.6         0.8         2.3         3.0           Localized pain         2.0         1.6         0.9         0	Convulsion	0	0	2.8	2.0
Thrombocythemia         0         0         2.8         2.0           Cough         2.4         4.0         0.9         0           Generalized abdominal pain         2.4         2.8         0.9         2.0           Localized abdominal pain         2.4         2.8         0.5         1.0           Apnea         0         0         2.3         2.0           Gastrointestinal bleeding         0         0         2.3         1.0           Generalized edema         0         0         2.3         1.0           Loose stools         1.6         0.8         2.3         3.0           Localized pain         2.0         1.6         0.9         0	Hypokalemia	0	0	2.8	3.0
Cough         2.4         4.0         0.9         0           Generalized abdominal pain         2.4         2.8         0.9         2.0           Localized abdominal pain         2.4         2.8         0.5         1.0           Apnea         0         0         2.3         2.0           Gastrointestinal bleeding         0         0         2.3         1.0           Generalized edema         0         0         2.3         1.0           Loose stools         1.6         0.8         2.3         3.0           Localized pain         2.0         1.6         0.9         0	Pneumonia	0	0	2.8	2.0
Generalized abdominal pain         2.4         2.8         0.9         2.0           Localized abdominal pain         2.4         2.8         0.5         1.0           Apnea         0         0         2.3         2.0           Gastrointestinal bleeding         0         0         2.3         1.0           Generalized edema         0         0         2.3         1.0           Loose stools         1.6         0.8         2.3         3.0           Localized pain         2.0         1.6         0.9         0	Thrombocythemia	0	0	2.8	2.0
Localized abdominal pain         2.4         2.8         0.5         1.0           Apnea         0         0         2.3         2.0           Gastrointestinal bleeding         0         0         2.3         1.0           Generalized edema         0         0         2.3         1.0           Loose stools         1.6         0.8         2.3         3.0           Localized pain         2.0         1.6         0.9         0	Cough	2.4	4.0	0.9	0
Apnea         0         0         2.3         2.0           Gastrointestinal bleeding         0         0         2.3         1.0           Generalized edema         0         0         2.3         1.0           Loose stools         1.6         0.8         2.3         3.0           Localized pain         2.0         1.6         0.9         0	Generalized abdominal pain	2.4	2.8	0.9	2.0
Gastrointestinal bleeding         0         0         2.3         1.0           Generalized edema         0         0         2.3         1.0           Loose stools         1.6         0.8         2.3         3.0           Localized pain         2.0         1.6         0.9         0	Localized abdominal pain	2.4	2.8	0.5	1.0
Generalized edema         0         0         2.3         1.0           Loose stools         1.6         0.8         2.3         3.0           Localized pain         2.0         1.6         0.9         0	Apnea	0	0	2.3	2.0
Loose stools         1.6         0.8         2.3         3.0           Localized pain         2.0         1.6         0.9         0	Gastrointestinal bleeding	0	0	2.3	1.0
Localized pain 2.0 1.6 0.9 0	Generalized edema	0	0	2.3	1.0
1	Loose stools	1.6	0.8	2.3	3.0
Skin disorder         2.0         0         0.9         1.0	Localized pain	2.0	1.6	0.9	0
	Skin disorder	2.0	0	0.9	1.0

<sup>\*</sup> Patients 5 through 11 years of age received ZYVOX 10 mg/kg PO q12h or cefadroxil 15 mg/kg PO q12h. Patients 12 years or older received ZYVOX 600 mg PO q12h or cefadroxil 500 mg PO q12h.

Table 9 shows the incidence of drug-related adverse events reported in more than 1% of pediatric patients (and more than 1 patient) in either treatment group in the comparator-controlled Phase 3 trials.

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<sup>&</sup>lt;sup>†</sup> Patients from birth through 11 years of age received ZYVOX 10 mg/kg IV/PO q8h or vancomycin 10 to 15 mg/kg IV q6-24h, depending on age and renal clearance.

Table 9. Incidence (%) of Drug-related Adverse Events Occurring in >1% of Pediatric Patients (and >1 Patient) in Either Treatment Group in Comparator-Controlled Clinical Trials

		Skin and Skin Infections*	All Other Indications <sup>†</sup>		
Event	ZYVOX (n=248)	Cefadroxil (n=251)	ZYVOX (n=215)	Vancomycin (n=101)	
% of patients with ≥1 drug- related adverse event	19.2	14.1	18.8	34.3	
% of patients discontinuing due to a drug-related adverse event	1.6	2.4	0.9	6.1	
Diarrhea	5.7	5.2	3.8	6.1	
Nausea	3.3	2.0	1.4	0	
Headache	2.4	0.8	0	0	
Loose stools	1.2	0.8	1.9	0	
Thrombocytopenia	0	0	1.9	0	
Vomiting	1.2	2.4	1.9	1.0	
Generalized abdominal pain	1.6	1.2	0	0	
Localized abdominal pain	1.6	1.2	0	0	
Anemia	0	0	1.4	1.0	
Eosinophilia	0.4	0.4	1.4	0	
Rash	0.4	1.2	1.4	7.1	
Vertigo	1.2	0.4	0	0	
Oral moniliasis	0	0	0.9	4.0	
Fever	0	0	0.5	3.0	
Pruritus at non-application site	0.4	0	0	2.0	
Anaphylaxis	0	0	0	10.1 <sup>‡</sup>	

<sup>\*</sup> Patients 5 through 11 years of age received ZYVOX 10 mg/kg PO q12h or cefadroxil 15 mg/kg PO q12h. Patients 12 years or older received ZYVOX 600 mg PO q12h or cefadroxil 500 mg PO q12h.

## **Laboratory Changes**

ZYVOX has been associated with thrombocytopenia when used in doses up to and including 600 mg every 12 hours for up to 28 days. In Phase 3 comparator-controlled trials, the percentage of adult patients who developed a substantially low platelet count (defined as less than 75% of lower limit of normal and/or baseline) was 2.4% (range among studies: 0.3 to 10.0%) with ZYVOX and 1.5% (range among studies: 0.4 to 7.0%) with a comparator. In a study of hospitalized pediatric patients ranging in age from birth through 11 years, the percentage of patients who developed a substantially low platelet count (defined as less than 75% of lower limit of normal and/or baseline) was 12.9% with ZYVOX and 13.4% with vancomycin. In an outpatient study of pediatric patients aged from 5 through 17 years, the percentage of patients who developed a substantially low platelet count was 0% with ZYVOX and 0.4% with cefadroxil. Thrombocytopenia associated with the use of ZYVOX appears to be dependent on duration of therapy, (generally greater than 2 weeks of treatment). The platelet counts for most patients returned to the normal range/baseline during the follow-up period. No related clinical adverse events were identified in Phase 3 clinical trials in patients developing thrombocytopenia. Bleeding events were identified in thrombocytopenic patients in a

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Patients from birth through 11 years of age received ZYVOX 10 mg/kg IV/PO q8h or vancomycin 10 to 15 mg/kg IV q6-24h, depending on age and renal clearance.

<sup>&</sup>lt;sup>‡</sup> These reports were of 'red-man syndrome', which were coded as anaphylaxis.

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786 787 compassionate use program for ZYVOX; the role of linezolid in these events cannot be determined (see **WARNINGS**).

Changes seen in other laboratory parameters, without regard to drug relationship, revealed no substantial differences between ZYVOX and the comparators. These changes were generally not clinically significant, did not lead to discontinuation of therapy, and were reversible. The incidence of adult and pediatric patients with at least one substantially abnormal hematologic or serum chemistry value is presented in Tables 10, 11, 12, and 13.

Table 10. Percent of Adult Patients who Experienced at Least One Substantially Abnormal\* Hematology Laboratory Value in Comparator-Controlled Clinical Trials with ZYVOX

	Thats with 21 vozt						
Laboratory Assay	-	l Skin and Skin Infections	All Other Indications				
	ZYVOX Clarithromycin 400 mg q12h 250 mg q12h		ZYVOX 600 mg q12h	All Other Comparators <sup>†</sup>			
Hemoglobin (g/dL)	0.9	0.0	7.1	6.6			
Platelet count (x 10 <sup>3</sup> /mm <sup>3</sup> )	0.7	0.8	3.0	1.8			
WBC (x $10^{3}/\text{mm}^{3}$ )	0.2	0.6	2.2	1.3			
Neutrophils (x 10 <sup>3</sup> /mm <sup>3</sup> )	0.0	0.2	1.1	1.2			

<sup>&</sup>lt;75% (<50% for neutrophils) of Lower Limit of Normal (LLN) for values normal at baseline; <75% (<50% for neutrophils) of LLN and of baseline for values abnormal at baseline.

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Table 11. Percent of Adult Patients who Experienced at Least One Substantially Abnormal\* Serum Chemistry Laboratory Value in Comparator-Controlled Clinical Trials with ZYVOX

Laboratory Assay	_	d Skin and Skin Infections	All Other Indications		
	ZYVOX 400 mg q12h	Clarithromycin 250 mg q12h	ZYVOX 600 mg q12h	All Other Comparators <sup>†</sup>	
AST (U/L)	1.7	1.3	5.0	6.8	
ALT (U/L)	1.7	1.7	9.6	9.3	
LDH (U/L)	0.2	0.2	1.8	1.5	
Alkaline phosphatase (U/L)	0.2	0.2	3.5	3.1	
Lipase (U/L)	2.8	2.6	4.3	4.2	
Amylase (U/L)	0.2	0.2	2.4	2.0	
Total bilirubin (mg/dL)	0.2	0.0	0.9	1.1	
BUN (mg/dL)	0.2	0.0	2.1	1.5	
Creatinine (mg/dL)	0.2	0.0	0.2	0.6	

<sup>&</sup>gt;2 x Upper Limit of Normal (ULN) for values normal at baseline;

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<sup>&</sup>lt;sup>†</sup> Comparators included cefpodoxime proxetil 200 mg PO q12h; ceftriaxone 1 g IV q12h; dicloxacillin 500 mg PO q6h; oxacillin 2 g IV q6h; vancomycin 1 g IV q12h.

<sup>&</sup>gt;2 x ULN and >2 x baseline for values abnormal at baseline.

<sup>&</sup>lt;sup>†</sup> Comparators included cefpodoxime proxetil 200 mg PO q12h; ceftriaxone 1 g IV q12h; dicloxacillin 500 mg PO q6h; oxacillin 2 g IV q6h; vancomycin 1 g IV q12h.

Table 12. Percent of Pediatric Patients who Experienced at Least One Substantially Abnormal\* Hematology Laboratory Value in Comparator-Controlled Clinical Trials with ZYVOX

		LIVOA			
Laboratory Assay	Uncomplicated Skin and Skin Structure Infections <sup>†</sup>		All Other Indications <sup>‡</sup>		
	ZYVOX	Cefadroxil	ZYVOX	Vancomycin	
Hemoglobin (g/dL)	0.0	0.0	15.7	12.4	
Platelet count (x 10 <sup>3</sup> /mm <sup>3</sup> )	0.0	0.4	12.9	13.4	
WBC (x $10^{3}/\text{mm}^{3}$ )	0.8	0.8	12.4	10.3	
Neutrophils (x 10 <sup>3</sup> /mm <sup>3</sup> )	1.2	0.8	5.9	4.3	

<sup>&</sup>lt;75% (<50% for neutrophils) of Lower Limit of Normal (LLN) for values normal at baseline;

Table 13. Percent of Pediatric Patients who Experienced at Least One Substantially Abnormal\* Serum Chemistry Laboratory Value in Comparator-Controlled Clinical Trials with ZYVOX

<b>Laboratory Assay</b>	Uncomplicated Skin and Skin Structure Infections <sup>†</sup>		All Other Indications <sup>‡</sup>		
	ZYVOX	Cefadroxil	ZYVOX	Vancomycin	
ALT (U/L)	0.0	0.0	10.1	12.5	
Lipase (U/L)	0.4	1.2			
Amylase (U/L)			0.6	1.3	
Total bilirubin (mg/dL)			6.3	5.2	
Creatinine (mg/dL)	0.4	0.0	2.4	1.0	

<sup>&</sup>gt;2 x Upper Limit of Normal (ULN) for values normal at baseline; >2 x ULN and >2 (>1.5 for total bilirubin) x baseline for values abnormal at baseline.

## **Postmarketing Experience**

Myelosuppression (including anemia, leukopenia, pancytopenia, and thrombocytopenia) has been reported during postmarketing use of ZYVOX (see WARNINGS). Peripheral neuropathy, and optic neuropathy sometimes progressing to loss of vision, have been reported in patients treated with ZYVOX. Lactic acidosis has been reported with the use of ZYVOX (see PRECAUTIONS, General). Although these reports have primarily been in patients treated for longer than the maximum recommended duration of 28 days, these events have also been reported in patients receiving shorter courses of therapy. Serotonin syndrome has been reported in patients receiving concomitant serotonergic agents, including antidepressants such as selective serotonin reuptake inhibitors (SSRIs) and ZYVOX (see PRECAUTIONS, General). Convulsions have been reported with the use of ZYVOX (see PRECAUTIONS, General). Anaphylaxis, angioedema, and bullous skin

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<sup>&</sup>lt;75% (<50% for neutrophils) of LLN and <75% (<50% for neutrophils, <90% for hemoglobin if baseline <LLN) of baseline for values abnormal at baseline.

<sup>&</sup>lt;sup>†</sup> Patients 5 through 11 years of age received ZYVOX 10 mg/kg PO q12h or cefadroxil 15 mg/kg PO q12h. Patients 12 years or older received ZYVOX 600 mg PO q12h or cefadroxil 500 mg PO q12h.

Patients from birth through 11 years of age received ZYVOX 10 mg/kg IV/PO q8h or vancomycin 10 to 15 mg/kg IV q6-24h, depending on age and renal clearance.

<sup>&</sup>lt;sup>†</sup> Patients 5 through 11 years of age received ZYVOX 10 mg/kg PO q12h or cefadroxil 15 mg/kg PO q12h. Patients 12 years or older received ZYVOX 600 mg PO q12h or cefadroxil 500 mg PO q12h.

<sup>&</sup>lt;sup>‡</sup> Patients from birth through 11 years of age received ZYVOX 10 mg/kg IV/PO q8h or vancomycin 10 to 15 mg/kg IV q6-24h, depending on age and renal clearance.

- 807 disorders such as those described as Stevens Johnson syndrome have been reported. 808 Superficial tooth discoloration and tongue discoloration have been reported with the use of linezolid. The tooth discoloration was removable with professional dental cleaning (manual 809 descaling) in cases with known outcome. Hypoglycemia, including symptomatic episodes, 810 has been reported (see **WARNINGS**). These events have been chosen for inclusion due to 811 either their seriousness, frequency of reporting, possible causal connection to ZYVOX, or a 812 combination of these factors. Because they are reported voluntarily from a population of 813 unknown size, estimates of frequency cannot be made and causal relationship cannot be 814 815 precisely established.
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#### OVERDOSAGE

- In the event of overdosage, supportive care is advised, with maintenance of glomerular
- filtration. Hemodialysis may facilitate more rapid elimination of linezolid. In a Phase 1
- 820 clinical trial, approximately 30% of a dose of linezolid was removed during a 3-hour
- hemodialysis session beginning 3 hours after the dose of linezolid was administered. Data
- are not available for removal of linezolid with peritoneal dialysis or hemoperfusion.
- 823 Clinical signs of acute toxicity in animals were decreased activity and ataxia in rats and
- vomiting and tremors in dogs treated with 3000 mg/kg/day and 2000 mg/kg/day,
- respectively.
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#### DOSAGE AND ADMINISTRATION

- The recommended dosage for ZYVOX formulations for the treatment of infections is described in Table 14.
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**Table 14. Dosage Guidelines for ZYVOX** 

	Dosage and Rou	te of Administration	Recommended
Infection*	Pediatric Patients <sup>†</sup> (Birth through 11 Years of Age)	Adults and Adolescents (12 Years and Older)	Duration of Treatment (consecutive days)
Complicated skin and skin structure infections Community-acquired pneumonia, including concurrent bacteremia Nosocomial pneumonia	10 mg/kg IV or oral <sup>‡</sup> q8h	600 mg IV or oral <sup>‡</sup> q12h	10 to 14
Vancomycin-resistant Enterococcus faecium infections, including concurrent bacteremia	10 mg/kg IV or oral <sup>‡</sup> q8h	600 mg IV or oral <sup>‡</sup> q12h	14 to 28
Uncomplicated skin and skin structure infections	<5 yrs: 10 mg/kg oral <sup>‡</sup> q8h 5-11 yrs: 10 mg/kg oral <sup>‡</sup> q12h	Adults: 400 mg oral <sup>‡</sup> q12h Adolescents: 600 mg oral <sup>‡</sup> q12h	10 to 14

<sup>\*</sup> Due to the designated pathogens (see **INDICATIONS AND USAGE**)

Adult patients with infection due to MRSA should be treated with ZYVOX 600 mg q12h.

In limited clinical experience, 5 out of 6 (83%) pediatric patients with infections due to Gram-positive pathogens with MICs of 4 µg/mL treated with ZYVOX had clinical cures. However, pediatric patients exhibit wider variability in linezolid clearance and systemic exposure (AUC) compared with adults. In pediatric patients with a sub-optimal clinical response, particularly those with pathogens with MIC of 4 µg/mL, lower systemic exposure, site and severity of infection, and the underlying medical condition should be considered when

assessing clinical response (see CLINICAL PHARMACOLOGY, Special Populations,

Pediatric and PRECAUTIONS, Pediatric Use).

In controlled clinical trials, the protocol-defined duration of treatment for all infections ranged from 7 to 28 days. Total treatment duration was determined by the treating physician based on site and severity of the infection, and on the patient's clinical response.

No dose adjustment is necessary when switching from intravenous to oral administration. Patients whose therapy is started with ZYVOX I.V. Injection may be switched to either ZYVOX Tablets or Oral Suspension at the discretion of the physician, when clinically indicated.

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Neonates <7 days: Most pre-term neonates < 7 days of age (gestational age < 34 weeks) have lower systemic linezolid clearance values and larger AUC values than many full-term neonates and older infants. These neonates should be initiated with a dosing regimen of 10 mg/kg q12h. Consideration may be given to the use of 10 mg/kg q8h regimen in neonates with a sub-optimal clinical response. All neonatal patients should receive 10 mg/kg q8h by 7 days of life (see CLINICAL PHARMACOLOGY, Special Populations, Pediatric).

<sup>&</sup>lt;sup>‡</sup> Oral dosing using either ZYVOX Tablets or ZYVOX for Oral Suspension

#### 853 Intravenous Administration

- 854 ZYVOX I.V. Injection is supplied in single-use, ready-to-use infusion bags (see **HOW**
- 855 **SUPPLIED** for container sizes). Parenteral drug products should be inspected visually for
- particulate matter prior to administration. Check for minute leaks by firmly squeezing the
- bag. If leaks are detected, discard the solution, as sterility may be impaired.

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- 24 ZYVOX I.V. Injection should be administered by intravenous infusion over a period of 30
- to 120 minutes. **Do not use this intravenous infusion bag in series connections**.
- Additives should not be introduced into this solution. If ZYVOX I.V. Injection is to be
- given concomitantly with another drug, each drug should be given separately in accordance
- with the recommended dosage and route of administration for each product. In particular,
- physical incompatibilities resulted when ZYVOX I.V. Injection was combined with the
- 865 following drugs during simulated Y-site administration: amphotericin B, chlorpromazine
- 866 HCl, diazepam, pentamidine isothionate, erythromycin lactobionate, phenytoin sodium, and
- trimethoprim-sulfamethoxazole. Additionally, chemical incompatibility resulted when
- 868 ZYVOX I.V. Injection was combined with ceftriaxone sodium.

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- 870 If the same intravenous line is used for sequential infusion of several drugs, the line should
- be flushed before and after infusion of ZYVOX I.V. Injection with an infusion solution
- compatible with ZYVOX I.V. Injection and with any other drug(s) administered via this
- common line (see Compatible Intravenous Solutions).

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## **Compatible Intravenous Solutions**

- 876 5% Dextrose Injection, USP
- 877 0.9% Sodium Chloride Injection, USP
- 878 Lactated Ringer's Injection, USP

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- 880 Keep the infusion bags in the overwrap until ready to use. Store at room temperature.
- Protect from freezing. ZYVOX I.V. Injection may exhibit a yellow color that can intensify
- over time without adversely affecting potency.

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## **Constitution of Oral Suspension**

- 24 ZYVOX for Oral Suspension is supplied as a powder/granule for constitution. Gently tap
- bottle to loosen powder. Add a total of 123 mL distilled water in two portions. After
- adding the first half, shake vigorously to wet all of the powder. Then add the second half of
- the water and shake vigorously to obtain a uniform suspension. After constitution, each 5
- mL of the suspension contains 100 mg of linezolid. Before using, gently mix by inverting
- the bottle 3 to 5 times. **DO NOT SHAKE.** Store constituted suspension at room
- temperature. Use within 21 days after constitution.

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#### HOW SUPPLIED

- Injection
- 895 ZYVOX I.V. Injection is available in single-use, ready-to-use flexible plastic infusion bags
- in a foil laminate overwrap. The infusion bags and ports are latex-free. The infusion bags
- are available in the following package sizes:

- 899 100 mL bag (200 mg linezolid)
- NDC 0009-5137-01

900       200 mL bag (400 mg linezolid)       NDC 0009-5139-01         901       300 mL bag (600 mg linezolid)       NDC 0009-5140-01         902       Tablets         904       ZYVOX Tablets are available as follows:         905       400 mg (white, oblong, film-coated tablets printed with "ZYVOX 400mg")         907       100 tablets in HDPE bottle       NDC 0009-5134-01         908       20 tablets in HDPE bottle       NDC 0009-5134-02         909       Unit dose packages of 30 tablets       NDC 0009-5134-03	
902 903 <b>Tablets</b> 904 ZYVOX Tablets are available as follows: 905 906 <b>400 mg</b> (white, oblong, film-coated tablets printed with "ZYVOX 400mg") 907 100 tablets in HDPE bottle NDC 0009-5134-01 908 20 tablets in HDPE bottle NDC 0009-5134-02 909 Unit dose packages of 30 tablets NDC 0009-5134-03	
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905 906 400 mg (white, oblong, film-coated tablets printed with "ZYVOX 400mg") 907 100 tablets in HDPE bottle NDC 0009-5134-01 908 20 tablets in HDPE bottle NDC 0009-5134-02 909 Unit dose packages of 30 tablets NDC 0009-5134-03	
<ul> <li>400 mg (white, oblong, film-coated tablets printed with "ZYVOX 400mg")</li> <li>100 tablets in HDPE bottle</li> <li>20 tablets in HDPE bottle</li> <li>NDC 0009-5134-01</li> <li>NDC 0009-5134-02</li> <li>Unit dose packages of 30 tablets</li> <li>NDC 0009-5134-03</li> </ul>	
<ul> <li>400 mg (white, oblong, film-coated tablets printed with "ZYVOX 400mg")</li> <li>100 tablets in HDPE bottle</li> <li>20 tablets in HDPE bottle</li> <li>NDC 0009-5134-01</li> <li>NDC 0009-5134-02</li> <li>Unit dose packages of 30 tablets</li> <li>NDC 0009-5134-03</li> </ul>	
907       100 tablets in HDPE bottle       NDC 0009-5134-01         908       20 tablets in HDPE bottle       NDC 0009-5134-02         909       Unit dose packages of 30 tablets       NDC 0009-5134-03	
908 20 tablets in HDPE bottle NDC 0009-5134-02 909 Unit dose packages of 30 tablets NDC 0009-5134-03	
909 Unit dose packages of 30 tablets NDC 0009-5134-03	
910	
911 <b>600 mg</b> (white, capsule-shaped, film-coated tablets printed with "ZYVOX 600 mg")	
912 100 tablets in HDPE bottle NDC 0009-5135-01	
913 20 tablets in HDPE bottle NDC 0009-5135-01	
914 Unit dose packages of 30 tablets NDC 0009-5135-02	
914 Offit dose packages of 30 tablets NDC 0009-3133-03	
916 Oral Suspension	
217 ZYVOX for Oral Suspension is available as a dry, white to off-white, orange-flavored	
granule/powder. When constituted as directed, each bottle will contain 150 mL of a suspension providing the equivalent of 100 mg of linezolid per each 5 mL. ZYVOX for	
920 Oral Suspension is supplied as follows:	r
921 222 100 m = /5 m L in 240 m L alara hadda NDC 0000 5126 01	rΓ
922 100 mg/5 mL in 240-mL glass bottles NDC 0009-5136-01	r
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923 924 Storage of ZYVOX Formulations 925 Store at 25°C (77°F); excursions permitted to 15-30°C (59-86°F) [see USP Controlled Roor 926 Temperature]. Protect from light. Keep bottles tightly closed to protect from moisture. It 927 is recommended that the infusion bags be kept in the overwrap until ready to use. Protect 928 infusion bags from freezing. 929 930 CLINICAL STUDIES 931 932 Adults 933 Vancomycin-Resistant Enterococcal Infections 934 Adult patients with documented or suspected vancomycin-resistant enterococcal infection 935 were enrolled in a randomized, multi-center, double-blind trial comparing a high dose of 936 ZYVOX (600 mg) with a low dose of ZYVOX (200 mg) given every 12 hours (q12h) eithe 937 intravenously (IV) or orally for 7 to 28 days. Patients could receive concomitant aztreonan 938 or aminoglycosides. There were 79 patients randomized to high-dose linezolid and 66 to 939 low-dose linezolid. The intent-to-treat (ITT) population with documented vancomycin- 940 resistant enterococcal infection at baseline consisted of 65 patients in the high-dose arm an 941 52 in the low-dose arm.	Room It lect  ion of either onam to - n and
923 924 Storage of ZYVOX Formulations 925 Store at 25°C (77°F); excursions permitted to 15-30°C (59-86°F) [see USP Controlled Roor 926 Temperature]. Protect from light. Keep bottles tightly closed to protect from moisture. It 927 is recommended that the infusion bags be kept in the overwrap until ready to use. Protect 928 infusion bags from freezing. 929 930 CLINICAL STUDIES 931 932 Adults 933 Vancomycin-Resistant Enterococcal Infections 934 Adult patients with documented or suspected vancomycin-resistant enterococcal infection 935 were enrolled in a randomized, multi-center, double-blind trial comparing a high dose of 936 ZYVOX (600 mg) with a low dose of ZYVOX (200 mg) given every 12 hours (q12h) eithe 937 intravenously (IV) or orally for 7 to 28 days. Patients could receive concomitant aztreonan 938 or aminoglycosides. There were 79 patients randomized to high-dose linezolid and 66 to 939 low-dose linezolid. The intent-to-treat (ITT) population with documented vancomycin- 940 resistant enterococcal infection at baseline consisted of 65 patients in the high-dose arm an 941 52 in the low-dose arm. 942 943 The cure rates for the ITT population with documented vancomycin-resistant enterococcal	Room It lect  ion of either onam to - n and
923 924 Storage of ZYVOX Formulations 925 Store at 25°C (77°F); excursions permitted to 15-30°C (59-86°F) [see USP Controlled Roor 926 Temperature]. Protect from light. Keep bottles tightly closed to protect from moisture. It 927 is recommended that the infusion bags be kept in the overwrap until ready to use. Protect 928 infusion bags from freezing. 929 930 CLINICAL STUDIES 931 932 Adults 933 Vancomycin-Resistant Enterococcal Infections 934 Adult patients with documented or suspected vancomycin-resistant enterococcal infection 935 were enrolled in a randomized, multi-center, double-blind trial comparing a high dose of 936 ZYVOX (600 mg) with a low dose of ZYVOX (200 mg) given every 12 hours (q12h) eithe 937 intravenously (IV) or orally for 7 to 28 days. Patients could receive concomitant aztreonan 938 or aminoglycosides. There were 79 patients randomized to high-dose linezolid and 66 to 939 low-dose linezolid. The intent-to-treat (ITT) population with documented vancomycin- 940 resistant enterococcal infection at baseline consisted of 65 patients in the high-dose arm an 941 52 in the low-dose arm.	Room It lect  ion of either onam to - n and ecal do

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the high-dose arm than in the low-dose arm, although the difference was not statistically significant at the 0.05 level.

Table 15. Cure Rates at the Test-of-Cure Visit for ITT Adult Patients with Documented Vancomycin-Resistant Enterococcal Infections at Baseline

	Cu	red
Source of Infection	ZYVOX 600 mg q12h n/N (%)	ZYVOX 200 mg q12h n/N (%)
Any site	39/58 (67)	24/46 (52)
Any site with associated bacteremia	10/17 (59)	4/14 (29)
Bacteremia of unknown origin	5/10 (50)	2/7 (29)
Skin and skin structure	9/13 (69)	5/5 (100)
Urinary tract	12/19 (63)	12/20 (60)
Pneumonia	2/3 (67)	0/1 (0)
Other*	11/13 (85)	5/13 (39)

Includes sources of infection such as hepatic abscess, biliary sepsis, necrotic gall bladder, pericolonic abscess, pancreatitis, and catheter-related infection.

## **Nosocomial Pneumonia**

Adult patients with clinically and radiologically documented nosocomial pneumonia were enrolled in a randomized, multi-center, double-blind trial. Patients were treated for 7 to 21 days. One group received ZYVOX I.V. Injection 600 mg q12h, and the other group received vancomycin 1 g q12h IV. Both groups received concomitant aztreonam (1 to 2 g every 8 hours IV), which could be continued if clinically indicated. There were 203 linezolid-treated and 193 vancomycin-treated patients enrolled in the study. One hundred twenty-two (60%) linezolid-treated patients and 103 (53%) vancomycin-treated patients were clinically evaluable. The cure rates in clinically evaluable patients were 57% for linezolid-treated patients and 60% for vancomycin-treated patients. The cure rates in clinically evaluable patients with ventilator-associated pneumonia were 47% for linezolidtreated patients and 40% for vancomycin-treated patients. A modified intent-to-treat (MITT) analysis of 94 linezolid-treated patients and 83 vancomycin-treated patients included subjects who had a pathogen isolated before treatment. The cure rates in the MITT analysis were 57% in linezolid-treated patients and 46% in vancomycin-treated patients. The cure rates by pathogen for microbiologically evaluable patients are presented in Table 16.

Table 16. Cure Rates at the Test-of-Cure Visit for Microbiologically Evaluable Adult Patients with Nosocomial Pneumonia

	Cured		
Pathogen	ZYVOX n/N (%)	Vancomycin n/N (%)	
Staphylococcus aureus	23/38 (61)	14/23 (61)	
Methicillin-resistant S. aureus	13/22 (59)	7/10 (70)	
Streptococcus pneumoniae	9/9 (100)	9/10 (90)	

## Pneumonia caused by multi-drug resistant S.pneumoniae (MDRSP\*)

ZYVOX was studied for the treatment of community-acquired (CAP) and hospital-acquired (HAP) pneumonia due to MDRSP by pooling clinical data from seven comparative and non-comparative Phase 2 and Phase 3 studies involving adult and pediatric patients. The pooled MITT population consisted of all patients with *S.pneumoniae* isolated at baseline; the pooled ME population consisted of patients satisfying criteria for microbiologic evaluability. The pooled MITT population with CAP included 15 patients (41%) with severe illness (risk classes IV and V) as assessed by a prediction rule<sup>11</sup>. The pooled clinical cure rates for patients with CAP due to MDRSP were 35/48 (73%) in the MITT and 33/36 (92%) in the ME populations respectively. The pooled clinical cure rates for patients with HAP due to MDRSP were 12/18 (67%) in the MITT and 10/12 (83%) in the ME populations respectively.

Table 17. Clinical cure rates for 36 microbiologically-evaluable patients with CAP due to MDRSP\* who were treated with ZYVOX (stratified by antibiotic susceptibility)

Susceptibility Screening	Clinical Cure		
	n/N <sup>a</sup>	(%)	
Penicillin-resistant	14/16	88	
2 <sup>nd</sup> generation cephalosporin-resistant <sup>b</sup>	19/22	86	
Macrolide-resistant <sup>c</sup>	29/30	97	
Tetracycline-resistant	22/24	92	
Trimethoprim/sulfamethoxazole-resistant	18/21	86	

## **Complicated Skin and Skin Structure Infections**

Adult patients with clinically documented complicated skin and skin structure infections were enrolled in a randomized, multi-center, double-blind, double-dummy trial comparing study medications administered IV followed by medications given orally for a total of 10 to 21 days of treatment. One group of patients received ZYVOX I.V. Injection 600 mg q12h followed by ZYVOX Tablets 600 mg q12h; the other group received oxacillin 2 g every 6 hours (q6h) IV followed by dicloxacillin 500 mg q6h orally. Patients could receive concomitant aztreonam if clinically indicated. There were 400 linezolid-treated and 419 oxacillin-treated patients enrolled in the study. Two hundred forty-five (61%) linezolid-treated patients and 242 (58%) oxacillin-treated patients were elinically evaluable. The cure rates in clinically evaluable patients were 90% in linezolid-treated patients and 85% in oxacillin-treated patients. A modified intent-to-treat (MITT) analysis of 316 linezolid-treated patients and 313 oxacillin-treated patients included subjects who met all criteria for

a) n= pooled number of patients treated successfully; N= pooled number of patients having MDRSP isolates that exhibited resistance to the listed antibiotic

b) 2<sup>nd</sup>-generation cephalosporin tested was cefuroxime

c) macrolide tested was erythromycin

<sup>\*</sup> MDRSP refers to isolates resistant to two or more of the following antibiotics: penicillin, second-generation cephalosporins, macrolides, tetracycline, and trimethoprim/sulfamethoxazole.

study entry. The cure rates in the MITT analysis were 86% in linezolid-treated patients and 82% in oxacillin-treated patients. The cure rates by pathogen for microbiologically evaluable patients are presented in Table 18.

Table 18. Cure Rates at the Test-of-Cure Visit for Microbiologically Evaluable Adult Patients with Complicated Skin and Skin Structure Infections

		Cured
Pathogen	ZYVOX n/N (%)	Oxacillin/Dicloxacillin n/N (%)
Staphylococcus aureus	73/83 (88)	72/84 (86)
Methicillin-resistant <i>S. aureus</i>	2/3 (67)	0/0 (-)
Streptococcus agalactiae	6/6 (100)	3/6 (50)
Streptococcus pyogenes	18/26 (69)	21/28 (75)

A separate study provided additional experience with the use of ZYVOX in the treatment of methicillin-resistant *Staphylococcus aureus* (MRSA) infections. This was a randomized, open-label trial in hospitalized adult patients with documented or suspected MRSA infection.

One group of patients received ZYVOX I.V. Injection 600 mg q12h followed by ZYVOX Tablets 600 mg q12h. The other group of patients received vancomycin 1 g q12h IV. Both groups were treated for 7 to 28 days, and could receive concomitant aztreonam or gentamicin if clinically indicated. The cure rates in microbiologically evaluable patients with MRSA skin and skin structure infection were 26/33 (79%) for linezolid-treated patients and 24/33 (73%) for vancomycin-treated patients.

#### **Diabetic Foot Infections**

Adult diabetic patients with clinically documented complicated skin and skin structure infections ("diabetic foot infections") were enrolled in a randomized (2:1 ratio), multicenter, open-label trial comparing study medications administered IV or orally for a total of 14 to 28 days of treatment. One group of patients received ZYVOX 600 mg q12h IV or orally; the other group received ampicillin/sulbactam 1.5 to 3 g IV or amoxicillin/clavulanate 500 to 875 mg every 8 to 12 hours (q8-12h) orally. In countries where ampicillin/sulbactam is not marketed, amoxicillin/clavulanate 500 mg to 2 g every 6 hours (q6h) was used for the intravenous regimen. Patients in the comparator group could also be treated with vancomycin 1 g q12h IV if MRSA was isolated from the foot infection. Patients in either treatment group who had Gram-negative bacilli isolated from the infection site could also receive aztreonam 1 to 2 g q8-12h IV. All patients were eligible to receive appropriate adjunctive treatment methods, such as debridement and off-loading, as typically required in the treatment of diabetic foot infections, and most patients received these treatments. There were 241 linezolid-treated and 120 comparator-treated patients in the intent-to-treat (ITT) study population. Two hundred twelve (86%) linezolid-treated patients and 105 (85%) comparator-treated patients were clinically evaluable. In the ITT population, the cure rates were 68.5% (165/241) in linezolid-treated patients and 64% (77/120) in comparator-treated patients, where those with indeterminate and missing

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outcomes were considered failures. The cure rates in the clinically evaluable patients (excluding those with indeterminate and missing outcomes) were 83% (159/192) and 73% (74/101) in the linezolid- and comparator-treated patients, respectively. A critical post-hoc analysis focused on 121 linezolid-treated and 60 comparator-treated patients who had a Gram-positive pathogen isolated from the site of infection or from blood, who had less evidence of underlying osteomyelitis than the overall study population, and who did not receive prohibited antimicrobials. Based upon that analysis, the cure rates were 71% (86/121) in the linezolid-treated patients and 63% (38/60) in the comparator-treated patients. None of the above analyses were adjusted for the use of adjunctive therapies. The cure rates by pathogen for microbiologically evaluable patients are presented in Table 19.

Table 19. Cure Rates at the Test-of-Cure Visit for Microbiologically Evaluable Adult Patients with Diabetic Foot Infections

	Cı	ured
Pathogen	ZYVOX n/N (%)	Comparator n/N (%)
Staphylococcus aureus	49/63 (78)	20/29 (69)
Methicillin-resistant S. aureus	12/17 (71)	2/3 (67)
Streptococcus agalactiae	25/29 (86)	9/16 (56)

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#### **Pediatric Patients**

## **Infections Due to Gram-positive Organisms**

A safety and efficacy study provided experience on the use of ZYVOX in pediatric patients for the treatment of nosocomial pneumonia, complicated skin and skin structure infections, catheter-related bacteremia, bacteremia of unidentified source, and other infections due to Gram-positive bacterial pathogens, including methicillin-resistant and -susceptible Staphylococcus aureus and vancomycin-resistant Enterococcus faecium. Pediatric patients ranging in age from birth through 11 years with infections caused by the documented or suspected Gram-positive organisms were enrolled in a randomized, open-label, comparatorcontrolled trial. One group of patients received ZYVOX I.V. Injection 10 mg/kg every 8 hours (q8h) followed by ZYVOX for Oral Suspension 10 mg/kg q8h. A second group received vancomycin 10 to 15 mg/kg IV every 6 to 24 hours, depending on age and renal clearance. Patients who had confirmed VRE infections were placed in a third arm of the study and received ZYVOX 10 mg/kg q8h IV and/or orally. All patients were treated for a total of 10 to 28 days and could receive concomitant Gram-negative antibiotics if clinically indicated. In the intent-to-treat (ITT) population, there were 206 patients randomized to linezolid and 102 patients randomized to vancomycin. One hundred seventeen (57 %) linezolid-treated patients and 55 (54%) vancomycin-treated patients were clinically evaluable. The cure rates in ITT patients were 81% in patients randomized to linezolid and 83% in patients randomized to vancomycin (95% Confidence Interval of the treatment difference; -13%, 8%). The cure rates in clinically evaluable patients were 91% in linezolid-treated patients and 91% in vancomycin-treated patients (95% CI; -11%, 11%). Modified intent-to-treat (MITT) patients included ITT patients who, at baseline, had a Gram-positive pathogen isolated from the site of infection or from blood. The cure rates in MITT patients were 80% in patients randomized to linezolid and 90% in patients randomized to vancomycin (95% CI; -23%, 3%). The cure rates for ITT, MITT, and

clinically evaluable patients are presented in Table 20. After the study was completed, 13 additional patients ranging from 4 days through 16 years of age were enrolled in an open-label extension of the VRE arm of the study. Table 21 provides clinical cure rates by pathogen for microbiologically evaluable patients including microbiologically evaluable patients with vancomycin-resistant *Enterococcus faecium* from the extension of this study.

Table 20. Cure Rates at the Test-of-Cure Visit for Intent to Treat, Modified Intent to Treat, and Clinically Evaluable Pediatric Patients by Baseline Diagnosis

	IT	T	MI	$TT^*$	Clinically	Evaluable
Population	ZYVOX n/N (%)	Vancomycin n/N (%)	ZYVOX n/N (%)	Vancomycin n/N (%)	ZYVOX n/N (%)	Vancomycin n/N (%)
Any diagnosis	150/186 (81)	69/83 (83)	86/108 (80)	44/49 (90)	106/117 (91)	49/54 (91)
Bacteremia of unidentified source	22/29 (76)	11/16 (69)	8/12 (67)	7/8 (88)	14/17 (82)	7/9 (78)
Catheter-related bacteremia	30/41 (73)	8/12 (67)	25/35 (71)	7/10 (70)	21/25(84)	7/9 (78)
Complicated skin and skin structure infections	61/72 (85)	31/34 (91)	37/43 (86)	22/23 (96)	46/49 (94)	26/27 (96)
Nosocomial pneumonia	13/18 (72)	11/12 (92)	5/6 (83)	4/4 (100)	7/7 (100)	5/5 (100)
Other infections	24/26 (92)	8/9 (89)	11/12 (92)	4/4 (100)	18/19 (95)	4/4 (100)

MITT = ITT patients with an isolated Gram-positive pathogen at baseline

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Table 21. Cure Rates at the Test-of-Cure Visit for Microbiologically Evaluable Pediatric Patients with Infections due to Gram-positive Pathogens

	Microbiolog	ically Evaluable
Pathogen	ZYVOX n/N (%)	Vancomycin n/N (%)
Vancomycin-resistant Enterococcus faecium	6/8 (75)*	0/0 (-)
Staphylococcus aureus	36/38 (95)	23/24 (96)
Methicillin-resistant S. aureus	16/17 (94)	9/9 (100)
Streptococcus pyogenes	2/2 (100)	1/2 (50)

Includes data from 7 patients enrolled in the open-label extension of this study.

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